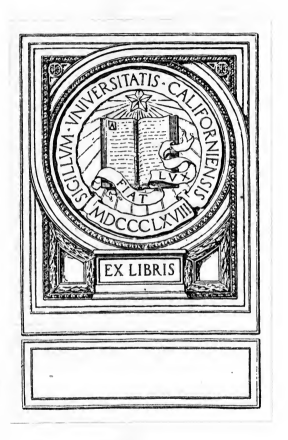
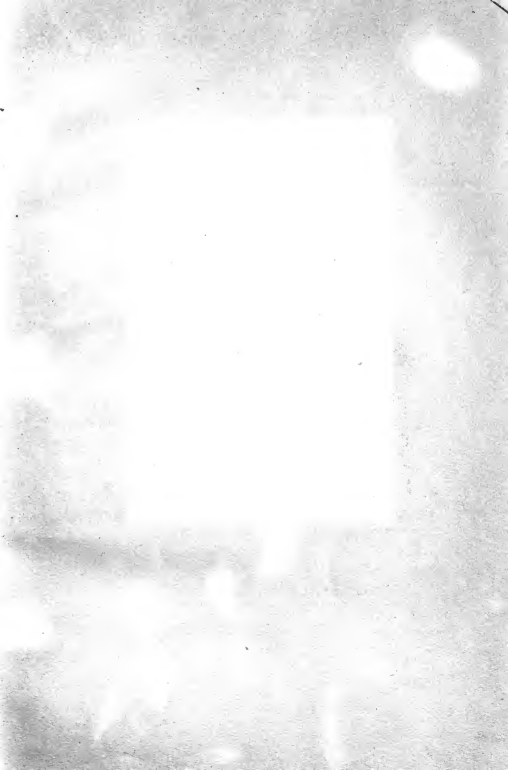
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ALEXANDER FINDLAY, M.A., D.Sc., F.I.C.

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MOLECULAR ASSOCIATION



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BY

W. E. S. TURNER, D.Sc. (London), M.Sc. (BIRMINGHAM)

LECTURER IN PHYSICAL AND INORGANIC CHEMISTRY IN THE UNIVERSITY
OF SHREFIELD

WITH DIAGRAMS



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PREFATORY NOTE.

Several of the author's students have taken an active interest in the production of this monograph. Mr. C. J. Peddle, M.Sc., has supplied the diagrams; Mr. C. T. Pollard, M.Sc., and in a very large measure Mr. C. C. Bissett, B.Sc., B.Met., have assisted in compiling the data contained in the Appendix; whilst the task of sorting and arranging these data has been carried out by Mr. S. English, M.Sc. To all these gentlemen the author expresses his hearty thanks.

The author will be glad to have his attention called by investigators to any work bearing on the subject of this monograph.



CONTENTS.

CHAPTER		PAGE
I.	Introductory	I
II.	Molecular Association in Gases. The molecular state of elementary gases and vapours, 10. Compounds, 12. The influence of temperature on molecular complexity, 14. The influence of pressure, 17.	6
III.	THE MOLECULAR COMPLEXITY OF DISSOLVED SUBSTANCES. The molecular state of the elements, 22. The molecular complexity of compounds, 25. Organic compounds, excluding salts, 26. Salts, 27. Solid solutions, 28. Colloidal solutions, 29. The effect of temperature on molecular weight, 31. The effect of concentration, 33.	21
IV.	THE MOLECULAR COMPLEXITY OF DISSOLVED SUBSTANCES. THE INFLUENCE OF THE SOLVENT	37
	influence of the solvent, 39. The effect of dielectric constant on electrolytes as solutes, 46.	
V.	Molecular Complexity in the Liquid State Evidence of molecular association in liquids; from vapour pressure data, 50. Boiling point evidence, 51. Latent heat, 51. Evidence from critical data, 53. Evidence from viscosity measurements, 55.	49
VI.	SURFACE TENSION METHODS OF DETERMINING MOLECULAR	
	COMPLEXITY IN THE LIQUID STATE	57

CONTENTS

CHAPTE		PAGI
VII.	Molecular Complexity in the Liquid State. Some other Methods and a Review. Longinescu's method, 74. Latent heat methods, 75. Bingham's fluidity method, 76. Traube's volume methods, 77. Guye's method, 79. Holmes' method, 81. General review of methods and results, 84.	74
VIII.	THE MOLECULAR COMPLEXITY OF WATER AND THE THEORY OF DYNAMIC ALLOTROPY	89
IX.	THE SELECTION AND USE OF MOLECULAR FORMULÆ	96
X.	MOLECULAR ASSOCIATION AND PHYSICAL PROPERTIES Degree of association and dielectric constant of a liquid, 100. Solvent power and degree of association, 101. Electrical conductivity and degree of association, 102. Molecular association and optical activity, 103.	100
XI.	Molecular Association and Chemical Combination	105
	References	110
	APPENDIX	119
	REFERENCES TO APPENDIX	165
	NDEX	160

CHAPTER I.

INTRODUCTORY.

In its widest sense, the term association embraces the whole of the phenomena of chemistry and various sections of physics. The smallest chemical unit, the atom, is made up, according to modern views, of associations of electrons the number and arrangement of which give the character to the individual atom; and by the association of these atoms, molecules composed either of similar or unlike atoms may be built up.

The subject of this monograph is concerned with the behaviour of the molecule, the dependence of its degree of complexity on the nature of the constituent atoms, and on the forces which act on it from without.

The molecule of an elementary substance may be of the simplest possible character like that of helium, monatomic; or, on the other hand, it may be of great complexity, like that of sulphur, which in the state of vapour near the boiling point of the substance, possesses on the average about eight atoms * in the molecule. Under the action of certain forces, however, such as heat, pressure, electrical stress, and solvent action, it is possible to reduce the size of the molecule, so that sulphur, for example, eventually even becomes monatomic. A substance such as sulphur at comparatively low temperatures has molecules which are associations of small ones, S_6 , S_9 , and S_1 , as will be shown later; it is, in short, an associated substance. The molecules of certain compound substances are likewise associated, such as those of aluminium chloride or arsenious oxide, and it is usually possible to split or dissociate such molecules into two or more like parts. Such a dissociation does not occur completely at some one point, but takes place gradu-

^{*} More correctly the vapour is a mixture of S₈ and S₆ molecules (see chap. 11.).

ally, so that over a considerable range a mixture of different kinds of molecules is obtained. This is true of all states of matter and all kinds of dissociation.

There is a second form of dissociation of a compound molecule which can take place whether the molecule is originally an association of smaller, like, or of unlike molecules. Nitrogen peroxide, for example, gradually dissociates with rise of temperature so that from one molecule NoO4, two like molecules NO9 are obtained. This process is complete at about 150°. Above 150°, a further dissociation sets in, when nitric oxide and oxygen, two unlike molecules, are produced, nitrogen peroxide being completely broken down into these products at 620° (under atmospheric pressure).1 The first type of dissociation may be called homogeneous, and it is with this type that the monograph is almost wholly concerned; the second, on the other hand, is heterogeneous dissociation. The relation of the two types, so far as is known, is the subject of one of the chapters of the book; but it may be pointed out here that the action of temperature and pressure in the two cases is similar.

Instances of molecular association in gas molecules were discovered quite early in the development of the atomic and molecular theories, as the result of vapour density determinations by Dumas and Mitscherlich, but were only clearly explained when Cannizzaro demonstrated how readily Avogadro's Hypothesis acted as a guide in the comparison of gas molecules. The general acceptance of the hypothesis naturally paved the way for the active investigation of the molecular state of gases and vapours, and towards this end, the invention of a simple form of apparatus by Viktor Meyer greatly assisted. From about 1880 onwards for some fifteen or sixteen years, Meyer and his co-workers, as well as Biltz, Nilson, and Pettersson, and others, who used either Meyer's or Dumas' apparatus, amassed a large amount of data relating to the vapour densities of metals, non-metals, oxides, and salts, data which were particularly valuable from the point of view of valency, the Periodic Table and particularly the variation of molecular weight with temperature. The form of the apparatus was gradually improved until temperatures up to about 1700°-1800° could both be attained and utilized. No considerable further advance was made, however, until Nernst introduced the method of electrical heating, and apparatus

of iridium, by means of which investigations were made possible up to temperatures exceeding 2000°. The extent to which our knowledge of the subject has followed the invention of new apparatus is again illustrated by the invention of the glass spiral manometer and its improved forms, which now enable densities of vapours at various temperatures and pressures to be measured with greatly increased accuracy.

While Meyer and others were actively engaged in the study of molecular weight at high temperatures, the investigations by Raoult² on the freezing points and vapour pressures of solutions provided a means of determining molecular weights at comparatively low temperatures; and the extension of the gas laws to solutions by van't Hoff provided a theoretical basis for Raoult's generalizations and set a seal on the great utility of freezing point and vapour pressure measurements. Two series of abnormal substances soon came to light. Thus, Raoult found that his generalizations could not be strictly applied to aqueous solutions of electrolytes, for the depression of the freezing point was always abnormally great—for binary electrolytes in dilute solution, about twice the value found with non-electrolytes. the other hand, a number of organic substances containing the hydroxyl group, when dissolved in solvents of the hydrocarbon type, depressed the freezing point less than was expected. According to the rules set out, therefore, it appeared as if a dissociation occurred in the first type, whilst in the second association took place, the molecular weight being greater than that corresponding to the simplest formula. To the second class, numerous examples of associated organic substances, alcohols, acids, and oximes were soon added as the result of the work of Paterno, Beckmann, Auwers, Fabyini, and Ampola and Manuelli. Recently, the behaviour of electrolytes has been studied much more completely and it has been shown that electrolytes are not only apparently dissociated in aqueous solutions, but also in certain other solvents, and that there is a gradual transition from dissociation to very marked association, according to the nature of the solvent. The molecular state in solution of elementary substances has likewise been studied by Raoult's processes, and it was in this country, quite early in the study of solutions, that the laws of Raoult were found to be equally valid for solutions of metals in metallic solvents; and

that, in consequence, the gas laws and Avogadro's Hypothesis could be applied to all types of solutions.

Methods for the investigation of pure liquids were only devised some years after it was possible to investigate solutions, although the properties of liquids had already been the subject of many investigations. To the question whether the size of molecules in liquids was the same as in the gaseous state. van der Waals, in 1876, replied that the continuous passage of a vapour into a liquid during which the relation between pressure and volume underwent no sharp change might be taken as indicating no alteration of molecular size. The first method of investigating molecular size was developed by Ramsay and Shields from previous deductions of Eötvös, and it was hoped that the method might be of very general application. Experience has shown that it breaks down outside a certain range of molecular weight. At the present time it must be admitted that although it is quite easy to detect association, the quantitative determination is another matter and there is often a big percentage difference between the results obtained by the many different processes of measurement that have been devised. There is great need of some guiding and reconciling principle like Avogadro's Hypothesis. With regard to the solid state, still less is known of molecular sizes, so little that no special section is called for in this monograph. The methods which have been proposed up to the present are extensions of those which yield results for liquids, and will be mentioned in the sections dealing with liquids.

The study of molecular complexity has had great influence on the progress of chemical theory. Reference has already been made to the density measurements of Dumas, made on mercury, iodine, sulphur, and phosphorus, and of Mitscherlich, on arsenic. Berzelius had previously, on the basis of Gay Lussac's Law of Volumes, believed that the relative densities of elementary gases were proportional to their atomic weights. Dumas and Mitscherlich showed that the atomic weights of mercury, sulphur, phosphorus, and arsenic so deduced were far different from the Berzelius' values found by chemical analysis; and Berzelius' atomic weight system falling into discredit, rival systems sprang up, none of which found general acceptance. Cannizzaro, introducing Avogadro's Hypothesis, explained the apparently con-

tradictory results with the elements named by giving a clear definition of the term molecule and of the relation of the atomic to the molecular weight. Different molecules, he demonstrated, had different degrees of complexity.

In the realm of solutions, too, the widely applied electrolytic dissociation hypothesis of Arrhenius attempted to explain the abnormal molecular weights of electrolytes in aqueous solution, whilst the phenomenon of osmosis has been explained by some chemists and physicists on the ground of association processes, both of like and unlike molecules.³

The newer theories of allotropy, too, are dependent on a study of molecular size. As a gas or vapour whose molecules can form aggregates contains, over a range of temperature, molecules of different sizes admixed, the proportions depending on the temperature and pressure, so also in liquids and solids such mixtures may exist. On this basis we have the theory of dynamic allotropy (chap. VIII.).

Finally we have chemical combination related in some way possibly to molecular association. This relationship forms the subject of the concluding chapter.

CHAPTER II.

MOLECULAR ASSOCIATION IN GASES.

In order to determine the molecular complexity of a gas, the molecular weight is found by means of a density measurement, and then compared with the simplest molecular weight obtained by chemical analysis; that is, compared with the sum of the atomic weights of the elements present. The method requires a knowledge of atomic weights, therefore, and for the elements which do not enter into chemical combination, recourse is had to a determination of the ratio, γ , of the specific heats of the gas or vapour, since for a monatomic gas $\gamma = 1.67$, for a diatomic gas 1.4 to 1.3, and decreases in value as the molecule becomes more complex. The determination of γ , indeed, serves to indicate in a few cases (monatomic vapours, for which alone it is trustworthy) the complexity independently of the density.

The density may be determined with accuracy by Regnault's method, approximately by Bunsen's. Whytlaw-Gray and Ramsay,⁴ for example, determined the density of niton by the former method; Debierne,⁵ for the same substance, used the latter. At high temperatures (2000°) Bunsen's method has received considerable application by Emich.⁶

More frequently employed in the study of molecular complexity, however, are the methods of Dumas,⁷ of Hofmann,⁸ and of V. Meyer,^{8a} particularly the first and last named, as both of them can be applied over a wide range of temperature. Of these two methods, that of Dumas can claim greater accuracy whilst the method of Meyer is more convenient.

The Dumas method, pretty much in its original form, has been made to yield results of fair precision by Biltz, but its most important development lies in the introduction of a device by which the pressure of the vapour within the bulb can be registered. The device consists of a spiral of glass or of quartz, or of a quartz membrane, or yet again, as in the original

suggestion of Ladenburg and Lehmann, of a flattened glass or quartz tube, the essential condition of any of the forms being its sensitiveness to pressure. The method ¹⁰ can be illustrated by reference to the diagram of the apparatus (Fig. I) employed by Preuner and Schupp, and to the following description.

The bulb A, in which the substance was vaporized, was made of quartz, and had an extension consisting of a flattened quartz

tube B in the form of a spiral, which ended in a quartz thread carrying the mirror. From the box or chamber C containing this mirror, connexion was made to a pump by which the external pressure could be altered and a mercury gauge by which the pressure was measured. Into the quartz bulb, the volume of which with the extension was known, a known weight of the material was introduced, and, after the air had been exhausted, the bulb was sealed at a, and with the spiral, heated in an electrical tube furnace to the temperature required, the actual temperature being recorded by a thermo-couple inserted in the tubulure at b. When the pressure within the bulb A, and therefore, also the spiral, rises, the spiral tends to unwind and in so doing the mirror is deflected by an amount which can be read from a scale by a microscope. On the other hand, when an external pressure acts on the spiral, the com-

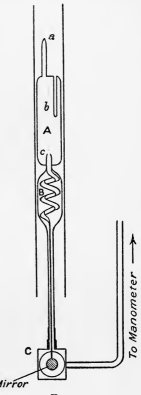


Fig. 1.

pression of the latter reverses the direction of the mirror deflection. Hence, in the actual determination, after the mirror had been deflected by the pressure of the vaporized substance, it was brought back to the zero point by the gradual application of an external pressure, this pressure, measured on the gauge, being equal to the vapour pressure of the substance. By this process, therefore, all the required factors, the weight of substance, its

volume, temperature and pressure, were obtained with considerable accuracy.*

Although by the preceding method density determinations can be carried out at temperatures up to 1200°, when quartz is employed, and in the simple Dumas' form at still higher temperatures with a porcelain bulb, Meyer's method undoubtedly has the advantage in work at high temperatures. The older investigators obtained constant temperatures by using the vapour from boiling liquids such as aniline (184°), diphenylamine (310°), sulphur (445°), phosphorus pentasulphide (518°), and stannous chloride (606°), and, for still higher temperatures, specially constructed gas furnaces. But electrical heating, obviously more capable of regulation than gas, has been used in the experiments of Nernst 11 and of von Wartenberg, 12 in which temperatures exceeding 2000° were attained. For the measurement of the temperature, Meyer 13 and his co-workers adopted the device of utilizing the vaporization bulb both for its original purpose and as an air thermometer. The introduction of the thermo-couple and of the optical pyrometer has afforded more convenient methods of temperature measurement, although Meyer's device which consisted in expelling the residual filling gas by hydrogen chloride or carbon dioxide into a tube over potash, and from the volume of the gas so expelled (the original volume of the bulb and the coefficients of expansion of the gas and of the material of the bulb being known) calculating the temperature attained, was capable of considerable accuracy.

Modern methods may be illustrated by reference to the apparatus of von Wartenberg (Fig. 2). Since porcelain softens at 1700° so much as to require protection, and platinum melts at 1755°, the vaporization bulb was made of iridium, its size being reduced to 3-4 c.c. capacity. The temperature required was obtained by electrically heating an iridium cylinder enclosed in a mass of magnesia, current being supplied by water-cooled copper leads. The stem of the vaporization apparatus had five connexions, A to the nitrogen or argon supply which was to fill the tube after air had been removed by a mercury pump connected through B; a tube C containing an arrangement, worked electro-magnetically, by which the stem could be closed at

^{*}For a discussion of the errors involved the original papers of Ladenburg and Lehmann, Johnson, Preuner, Starck and Bodenstein, and Gibson should be consulted.

will; a manometer DE and the storage tube F containing weighed amounts of the substance in tiny capsules of ignited magnesia which could be pushed into the vaporization apparatus by an iron rod actuated from without by a solenoid. As the bulb of the apparatus was small, the amount of substance had

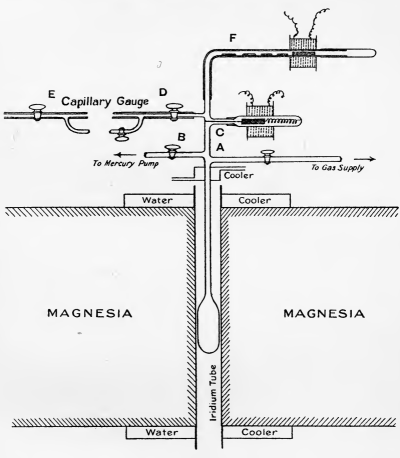


Fig. 2.

likewise to be reduced to a very small quantity, and the small expansion on vaporization was measured by the displacement in the calibrated capillary tube of the manometer of a drop of mercury moistened with sulphuric acid.

At high temperatures, metals become permeable to gases, and a glaze must be used as a protection. In von Wartenberg's

experiments, the bulb was coated within and without by a glaze of fused magnesia and magnesium chloride.

For the measurement of the temperature one of two methods was followed; either, an optical pyrometer was directed towards the bottom of the heated bulb, or, a rod of ceria and thoria was electrically heated until it glowed with a brightness equal to the bulb and the temperature of the rod measured optically.

The Molecular State of Gases and Vapours.

It will be convenient in giving an account of our present knowledge of the subject to quote first the results of measurements made on (a) elementary substances (b) compounds, and afterwards to deal more fully with the factors, namely, the temperature and the pressure influencing the molecular state.

Considering the elements as arranged in the groups of the Periodic Table, group O contains elements of the simplest molecular constitution. For the evidence of this we have to rely on determinations of the ratio of the specific heats from the velocity of sound. For helium, this ratio is 1.67,14 for argon 1.659,15 whilst the value for neon, xenon, and krypton is in each case approximately 1.66.16 As the ratio should be 1.67 for a monatomic and very approximately 1.4 for a diatomic gas, the suggestion has been made that argon contains a very small percentage of diatomic molecules. At present there is no other evidence to support this suggestion.

Of the elements in Group I, only four, hydrogen, sodium, potassium, and silver have received any study as gases. According to Longmuir, hydrogen is appreciably dissociated when the temperature exceeds 2100° (abs.). For trustworthy data on the alkali metals recourse must be had once again to the velocity of sound method, as all the density determinations appear to be open to criticism because of the corroding action of the metallic vapours. Recently, Wenz 19 has carried out a determination of the value of γ for potassium vapour at a temperature of 850°, using a steel tube silvered within. The value obtained, 1.77, although high, is sufficient evidence of the monatomicity of potassium vapour, and there is little doubt that the other alkali metals are similarly constituted in the state of vapour. In regard to silver, von Wartenberg's vapour density determinations

at about 2000° 20 likewise indicate the monatomic state of the molecule, although the results are not sufficiently accurate to decide whether molecules of a higher order also exist.

From Group II onwards, our knowledge of the molecular complexity of the elements has been gained by vapour density measurements. It may be summarized in the following table:—

MOLECULAR WEIGHTS BASED ON VAPOUR DENSITIES.

Element.	Periodic Group.	At. Weight.	Temperature.	Molecular Weight.
Silver 20	I	107.88	2000°	
Zinc 21	II	65.37	1400°	68.0-69.6
,, 22			1200°-2070°	66-76-7
Cadmium 23	,,	112'4	1700°	126 and 127.2
""		_	1 (111.0
Mercury 24	"	200.6	,	199.6
Thallium 25	111		1952°	201-205
Lead 22	IV	204	1320°-1690° 1600°	210-206
Leau	1 4	207.1	1870°	223
Phosphorus ²⁶	v	31.04	800°-1700°	210-239 111'3-92'1
Arsenic ²⁷		74.96	1700°	153.5-190.1
Antimony 22	,,	120*2	2070°	122-139
Bismuth 22 and 26	"	208.0	1640°	292
	- "	400 0	2070°	214-244
Sulphur ²⁸	VI	32.07	468°-606°	226-137
		3 . ,	1690°-2023°	59-54
Selenium 29	,,	79'2	1750°-1800°	178
,,			2070°	114-121
Tellurium 29	,,	127.5	1750°-1800°	264
,,,			2070°-2100°	152-179
Chlorine 30	VII	35•46	up to 1450°	normal
,,			>1450°	< 70.92
Bromine 31			o°-300°	>70.92
Bromine 31	,,	79.92	15°-750°	159.84
T 1' '90			800°-1050°	159.74-150.22
Iodine 32	,,	126.92	650°-1400°	254°-144°

The table is constructed from values determined at or near the atmospheric pressure. Determinations at various pressures and temperatures have also been made on phosphorus, arsenic, sulphur, and selenium and will be referred to in a later section; but it may be said here that all these elements have complex molecules which decrease in size with rise of temperature and diminution of pressure, until, in the case of sulphur and selenium, monatomic molecules are produced.

The results may be generalized in two ways. In the first place they may be regarded as illustrating once again the difference between the characteristics of metals and non-metals. It is true that the inert gases of the atmosphere resemble the metals in having molecules of the simplest character, namely monatomic, even at temperatures not far removed from their boiling points, but the other non-metallic elements are polyatomic, often diatomic, frequently, however, of a still higher order, so that at low temperatures the sulphur molecule contains eight atoms. There are but two substances with the properties of metals, antimony and bismuth, which have complex molecules, but the transition from the non-metallic to the metallic is nevertheless illustrated in the lower complexity of the bismuth molecule.

The results may also be reviewed from the standpoint of the Periodic Table. In the horizontal rows, elements to the left of Group V have so far as tests have been made monatomic molecules, elements in Groups V to VII polyatomic molecules, the maximum molecular complexity occurring in Group VI (sulphur and selenium). The ease with which a molecule can be split up depends also on the atomic weight of the element, as is illustrated by comparing arsenic, antimony, and bismuth, or, better still, by the halogen elements. Little is known of the molecular weight of fluorine, Moissan's 292 determination of the density giving a value less than that corresponding to F₂. Chlorine, however, between o° and 300° has an abnormally high vapour density, and the suggestion has been made that molecules Cl₄ 30a are present to the extent of 3:33 per cent at 0° and 0:02 per cent From this point up to 1450° the diatomic molecule is at 300°. stable.30 Bromine begins to split up into monatomic molecules at 750°-800° and iodine at a still lower temperature. Although molecular complexity tends to decrease in these cases with increase in mass, nitrogen and oxygen, first members of series of elements with complex molecules, are themselves only diatomic.

Compounds.

(a) Metallic.—

Substance.	Simplest Molecular Weight.	Observed Molecular Weight.	Temperature.
Silver chloride 13	143*3	165 239	1735° ? (lower than 1735)
Cuprous chloride 33	CuCl = 99.06	200	1560° 1700°
Glucinum chloride 84	80.06	186 88·6-80·7	589°-812°
Aluminium chloride 35		286.1-524.0	209°-440°
Aluminum emoride so	133*48	215.3-153.1	440°-1400°
Aluminium bromide 36	266°9	540'2	440°
Aluminium iodide 36	_	783.8	440°
Ferric chloride 33	407 ° 9 162 ° 2	321.0-318.5	447°-619°
Ferrous chloride 37	126.8	192.4-184.3	(yellow heat)
Stannous chloride 38	180.0	253.5-504.6	639°-1113°
Mercurous chloride 39	471	445	434.8°
	7/-	713	151

In addition to the substances tabulated, quite a number of other metallic chlorides have been investigated, NaCl, KCl, ¹¹ KI, ^{39a} ZnCl₂, ZnBr₂, CdBr₂, ³³ HgCl₂, GaCl₃, GaCl₂, InCl₃, InCl₂, InCl₅, ⁵¹ TlCl₂, PbCl₂, ⁵² ThCl₄, ⁵³ CrCl₃, CrCl₂, ⁵¹ but with the possible exception of InCl₂ and GaCl₃ there is no evidence of association. All the determinations save those of Nernst on sodium and potassium chlorides are not recent and might well be re-investigated by more accurate methods.

(b) Non-metallic Compounds .-

Substance.	Simplest Molecular Weight.	Observed Molecular Weight.	Temperature.
Water	18	(see p. 89)	26-4-88-1
Hydrogen fluoride ⁴⁰	20	51.6-20.7	
Hydrogen boride ⁴¹	?	B ₄ H ₁₀	
Hydrogen boride ⁴¹ Acetic acid ⁴² Metaphosphoric acid ⁴³ Nitrogen peroxide ⁴⁴ Phosphorous oxide ⁴⁵ Phosphoric oxide ⁴⁸	60.0 80.0 40.0	B ₆ H ₁₂ 100·6-70·8 154 and 156 76·6-40·7 224-225	120°-180° (? 1000°) 26·7°-154° 132°-184°
Phosphoric oxide ⁴³	142°1	370-307	(? 1000°)
Arsenious oxide ⁴⁶	197°9	396-199	518°-1800°
Antimony oxide ³³	288°4	566-577	1500°
Nitrogen trioxide ⁴⁷	76°02	76-2-119-6	ca.20°

The metallic compounds which are known to be associated in the state of vapour cannot very well be classified. They are all halogen compounds, it is true, but the obvious reason is that they alone, among the salts, can be volatilized without decomposition. It might be thought likely that the alkali haloids would be among the associated vapours, since, as will be seen later, they are associated in solution and in the fused state; but the existence of associated molecules in the vapour depends largely on the temperature coefficient of association. Liquid water has complex molecules which are rapidly broken up by rise of temperature, so that in the vapour, only a small proportion of double molecules are present, whilst the effect of temperature variation on acetic acid is much less and the vapour is far richer in double molecules. Despite the highly complex state of the fused alkali salts, therefore, it is not surprising that at the temperatures at which they may be volatilized, no appreciable association occurs.

Of the non-metallic associated vapours, most of them contain oxygen. Most of them, too, contain as the other element,

or one of the elements, a member of Group V. The analogous oxides, nitrogen trioxide, phosphorous, arsenious and antimonious oxides are all associated. The effect of temperature alteration on nitrogen trioxide has not yet been worked out. Phosphorous oxide undergoes heterogeneous dissociation 45 when heated, so that the only formula which will fit the substance is P_4O_6 . Mercurous chloride, of the metallic compounds, likewise yields $Hg + HgCl_2$, when heated. Although from Baker's work the likeliest formula is Hg_2Cl_2 , later work makes this doubtful. Cuprous chloride has not yet been broken down into CuCl molecules, nor have phosphoric oxide and metaphosphoric acid been split into smaller like molecules, so that their formulæ are Cu_2Cl_2 , P_4O_{10} and $H_2P_2O_6$ respectively.

The Influence of Temperature on Molecular Complexity.

From the very many examples already provided, the effect of high temperatures in simplifying molecular aggregations will be apparent. Just above its boiling point, sulphur molecules are so complex as to be represented on the average by the formula $S_7 - S_8$; but at 1000°, simplification to a molecule S_2 has occurred, whilst at 1000° higher still, the cleavage of diatomic into monatomic molecules is well on the way. This gradual dissociation of a complex molecule is well illustrated by Biltz's measurements on arsenious oxide at atmospheric pressure. In the following table, the values of the vapour density (d) compared with air, are given. From them the molecular weight, if desired, can be obtained by multiplying by the factor 28 9.

	DENSITY OF	ARSENIOUS OXIDE.	
· to.	d.	t°.	d_{\bullet}
518°	13.41-14.14	1450°	9.41
769°	13.62	1584°	8.80-8.83
851°	13.12	1732°	7'32
1059°	12.72-12.83	1800°	6.93
1256°	12.36		

Meyer believed that along the lines of high temperature measurement lay the hope of resolving the atoms of elements still further, and at the time of his death he had constructed a volatilization tube mainly of magnesia (88 per cent) for use at a temperature of 2000°. ⁴⁸ But even the conversion of diatomic into monatomic molecules is in a number of cases far from complete at this temperature, and no further division by heat has been accomplished in the case of a monatomic vapour, such as

mercury; the molecular weight of this substance at 1952° being, according to Nernst, still 201-205, the same value as at 360°.

The degree of dissociation can readily be calculated from the densities observed. Suppose, for example, that N molecules of the original complex form have been heated up to a temperature t^o , resulting in the dissociation of a certain fraction a (which fraction is the degree of dissociation), and that n simpler molecules result from the cleavage of each complex one. The number of molecules remaining undissociated will be N (I - a) and the number of simple molecules Nan. The total number of molecules, complex and simple, therefore, becomes

$$N(I - a) + Nan = N[I + a(n - I)].$$

With the increase in the number of molecules, the density diminishes in proportion. Hence, if d_o and d_t are the densities of the undissociated and partially dissociated vapours, then

$$\frac{d_t}{d_o} = \frac{N}{N \left[1 + a \left(n - 1\right)\right]} = \frac{1}{1 + a \left(n - 1\right)} \quad . \tag{1}$$
and
$$a = \frac{d_o - d_t}{d_t \left(n - 1\right)} \quad . \tag{2}$$

Since at every temperature the extent of dissociation has a definite value, the vapour of any dissociating substance consists of a mixture of different molecular forms. A reversible or equilibrium reaction thus exists, as in

$$\begin{split} & I_2 \rightleftarrows I + I \\ & N_2O_4 \rightleftarrows NO_2 + NO_2 \\ & (H_2O)_2 \rightleftarrows H_2O + H_2O \end{split}$$

and so on, and the law of mass action requires that the quantitative relationship

$$Kp = p_1p_2 = p_1^2$$
, shall hold,

where p measures the partial pressure and therefore the concentration of the still undissociated molecules, and p_1 and p_2 (which in all the above cases are equal), the corresponding partial pressures of the single molecules.

The manner in which a given complex molecule breaks down can therefore be tested by the validity of the equation expressing the equilibrium between the components of the mixture; for the value of K, the equilibrium constant, must remain unchanged at constant temperature. Thus, Preuner and Schupp ¹⁰ concluded that the dissociation of the sulphur molecule proceeded in the stages

gested by Erdmann 49.

(1)
$$3S_8 \rightleftharpoons 4S_6$$
, and (2) $S_6 \rightleftharpoons 3S_2$,

because the equilibrium equations $K_1 = \frac{p_1^4}{P_1^3}$ for the first and $K_2 = \frac{p_2^3}{P_2}$ for the second were found to be true; that is to say, the values of K_1 and K_2 were found to be constant. A third stage at high temperatures or low pressures also occurs, namely, $S_2 \gtrsim 2S$; but the expression $K_3 = \frac{p_3^4}{P_3}$, corresponding to a dissociation $S_8 \gtrsim 4S_2$, was not found to hold, and hence this mode of dissociation was ruled out as unlikely. Rejected also as improbable were the degrees of complexity S_7 and S_5 , whilst it was also impossible to reconcile the partial pressure measure-

In a very similar manner, Preuner and Brockmöller ¹⁰ came to the conclusion that the stages in the decomposition of selenium and arsenic were

ments with the existence of S3, a form analogous to O3 sug-

$$As_4 \rightarrow As_2 \rightarrow As_1$$

 $Se_6 \rightarrow Se_2 \rightarrow Se_1$

and Stock, Gibson, and Stamm similarly decided that between 500° and 1200° and at pressures ranging from 100 mm. to 1000 mm. the only change in the phosphorus molecule was represented by the transformation

$$P_4 \rightleftharpoons 2P_2$$

A study of the value of the equilibrium constant may furnish information on the energy changes accompanying a transformation from one kind of molecule to another. For the alteration in the point of equilibrium by a change of temperature is reflected in the value of the equilibrium constant, this alteration being expressed by the well-known equation of van't Hoff,

$$\frac{d \log_e K}{dT} = -\frac{Q}{RT^2},$$

T being the temperature measured on the absolute scale, R the gas constant and Q the amount of heat absorbed by a gram molecular quantity of substance during transformation. From the gas densities, the value of K can be obtained simply. For if the reaction be of the type

$$A_2 \gtrsim 2A$$
, then, as before,
 $Kp = p_1^2$.

The total pressure P of the gas mixture is the sum of the partial pressures $p + p_1$, and these in turn are proportional to the number of molecules of each form present. If, therefore, the degree of dissociation be a, the total number of molecules will increase in the proportion of N/N[1 + a(n-1)] (Equation 1), or 1: 1 + a since n = 2 in the case selected.

Hence,

$$p = P \frac{I - a}{I + a} = P \left(\frac{2d_t}{d} - I\right)$$
 from Equation 2,
and $p_1 = P \frac{2a}{I + a} = 2P \left(I - \frac{d}{d_0}\right)$.

Hence, also,

$$K = \frac{p_1^2}{p} = \frac{4 (d_o - d)^2 P}{(2d - d_o) d_o} . (3)^{50}$$

The value of Q, the latent heat of transformation, may now be calculated from the integrated form of the expression

$$\frac{d \log_e K}{dT} = -\frac{Q}{RT^2},$$

neglecting the small variation of Q with the temperature, namely,

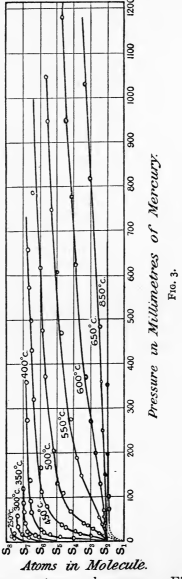
$$\log K = \frac{Q}{RT} + \text{constant}$$
 . (4).

In this way the following values have been obtained:—

The Influence of Pressure.

Although there are certain cases of dissociation (heterogeneous), such as $2HI \gtrsim H_2 + I_2$ on which pressure alteration has no effect because the number of molecules taking part in the direct and reverse reaction is the same, this never applies to homogeneous dissociation, which is under discussion in this chapter, as will be evident from the many examples quoted in

the preceding section. Recently most of the investigators of the dissociation of gas molecules have studied the simultaneous



effect of both temperature and pressure. Fig. 3 represents in graphic form Preuner and Schupp's results of the study of sulphur vapour under these conditions, and it will be seen that

even at 850° when the pressure is below 100 mm. the final stage in the dissociation is in progress.

Instead of utilizing these data to deduce the constants concerned, the influence of pressure on the dissociation of phosphorus vapour (Stock, Gibson, and Stamm) will be chosen for illustration. The value of the equilibrium constant is given by equation (3) and when known, it is possible to calculate the density of the substance at any temperature and pressure, since from

$$K = \frac{4 (d_o - d_t)^2 P}{(2d_t - d_o) d_o}, \text{ we obtain}$$

$$d_t = d_o - \frac{Kd_o}{4P} \left[\sqrt{1 + \frac{4P}{K} - 1} \right]^* \qquad (4)^{50}$$

In the following table are the values of K, of a, the degree of dissociation of the P_4 molecule, and of d_t observed and d_t calculated, using in the calculation, the mean value of K for the given temperature.

to.	α.	Р.	к.	dt (Obs.).	dt (Calcd.).
900°	0.02	606 mm.	6.06	58.7	5 8·9
,,	0.02	433 ,,	3.24	58.0	58.5
,,	0.02	243 ,,	4'32	57.8	57.5
,,	0'12	99 ,,	5.69	55.0	54.5
1200°	0.34	950 ,,	497	46.0	46.3
,,	0.38	699 ,,	461	44.8	44'7
,,,	0'48	412 ,,	489	4I°7	41.0
,,	0.01	175 ,,	413	38.3	37.8
]					

The values of the density are those relative to hydrogen as unit.

Not only may the density be re-calculated as indicated for the temperature for which K has been measured, but, with the assistance of equation (4) it can also be determined approximately for any other temperature and pressure, if the latent heat of transformation is known. The value of Q varies slightly with the temperature, but neglecting this, and inserting in the van't Hoff equation the value found at some lower temperature, K can be calculated for some higher or lower temperature. Thus, using the mean value of Q between 1100° and 1200°, namely,

$$d_t = d_o + \frac{K'}{P} - \frac{K'}{P} \sqrt{\frac{1 + \frac{d_o P}{K'}}{K'}}$$
 where $K' = \frac{Kd_o}{4}$.

^{*} This equation is more often employed in the form

- 49,200 cal., the value of K at 1300° was determined and from it in turn the degree of dissociation (of P_4) which should occur at different pressures. At 1300°, these values are:—

Р.	a.
I atmos.	60 per cent
0.5 "	69 ,, ,,
0.25 "	8r ", ",
100 mm.	89 ,, ,,

CHAPTER III.

THE MOLECULAR COMPLEXITY OF DISSOLVED SUBSTANCES.

THE extension, by van't Hoff, of the gas laws to solutions or to diluted matter in any form, makes it possible to determine the molecular weight of dissolved substances in a manner analogous to that employed for gases. The equation PV = RT applies in common. For gaseous substances, the molecular weight is that quantity which occupies the gram-molecular volume at 0° and 760 mm. pressure, and is determined by a density measurement. For solutions, on the other hand, it is customary to start with a known concentration (or density) and find the osmotic pressure corresponding to it. This osmotic pressure may be determined directly or indirectly. So far, the indirect methods, depending on the measurement of freezing point, or vapour pressure reduction or boiling point elevation, are the ones of most importance in practice.

Raoult's empirical laws 2 would, however, enable molecular weight determinations to be carried out independently of the van't Hoff, or osmotic pressure, theory. For although these laws rest in theory on the laws of osmotic pressure, and it is sometimes convenient to calculate the molecular depression (C) of freezing point or elevation of boiling point from van't Hoff's equation, $C = \frac{O \cdot O \cdot T^2}{I}$ where T is the absolute temperature and

I the latent heat of fusion or evaporation, yet in practice C is much more often determined by experiment, using some substance or substances generally accepted as normal. In this way the solvent employed is standardized.

Another method which is less often used is the partition coefficient method. If a substance is shaken with two immiscible solvents, each liquid appropriates a definite proportion of the whole such that if C_A is the concentration in solvent A and

 C_B that in solvent B, then $\frac{C_A}{C_B}$ is a constant, independent of the total amount dissolved, provided the molecular complexity is the same in each solvent. If, however, association or dissociation occurs in one solvent, or in both but to different extents, then $\frac{C_A}{C_B}$ is no longer constant; but if the solute is normal in A and associated n times the normal size in B, the ratio

$$\frac{C_A^n}{C_B}$$
 or $\frac{C_A}{\sqrt[n]{C_B}}$

is a constant quantity. From measurements of C_A and C_B , therefore, it is possible to determine the molecular complexity of a substance in one solvent relative to that in another. The process has been employed by Nernst, Hendrixson, Dawson, and Wedekind and Paschke.⁵⁴

As the practice of the various methods is so well known, the results will be at once discussed.⁵⁵

The Molecular State of the Elements.

(a) Metallic Elements.—Some of the earliest applications of the laws of Raoult were made with the view of extending them to metals. Thus we have the researches, in this field, of Heycock and Neville, Ramsay, and Tammann, 56 based, Ramsay's on vapour pressure, the others on freezing point measurements. The results obtained by Ramsay, who used mercury, and of Heycock and Neville who employed tin as solvent, are the more complete ones and will be the basis of consideration. The freezing point results are worked up so as to give, not the molecular weight, but the lowering produced by an atomic weight of the metal dissolved in 100 atoms of solvent.

Ramsay and Heycock and Neville came to the conclusion that most metals in metallic solvents have the simplest possible molecular constitution—monatomic. Heycock and Neville found that most of the metals produced much the same atomic lowering of freezing point in a given solvent, corresponding to Raoult's second law, and if, therefore, one metal could be regarded as monatomic then other metals, behaving similarly, must likewise be monatomic. This conclusion was very likely true because zinc and mercury in the state of vapour were known to be monatomic; and, further, from the van't Hoff

Solvent-Mercury.

Solvent-Tin.

VAPOUR PRESSURE RESULTS.

FREEZING POINT RESULTS.

Element.	At. Wt.	Atoms of element per 100 Atoms Hg.	Mol. Weight.	Atoms of element per 100 Atoms Sn.	Atomic Lowering.
Lithium Sodium Potassium Copper Silver Gold Calcium Barium Magnesium Zinc Cadmium Aluminium Gallium Indium Thallium Thallium Tin Lead Antimony Bismuth Manganese Nickel Palladium Mercury	6·9 23·0 39·1 63·6 107·9 197·2 40·1 137·4 24·3 65·4 112·4 207·1 69·9 114·8 204·0 119·0 207·1 20·2 08·0 154·9 258·7 166·7 200·6	1'70 0'86-5'35 1'55 and 5'26 3'22 1'59 and 2'80 2'9 0'90 0'70 and 4'82 1'07-3'16 0'75-3'51 1'52 and 2'94 2'27 0'81-3'33 1'94 and 3'94 2'00 and 4'39 1'86-6'22 3'77-4'68 1'14	7'1 21'6-15'1 29'1 and 30'2 112'4 207'4 and 208'1 19'1 75'7 24'0 and 21'5 70'1-65'4 100'2-103'8 36'7 and 33'1 69'7 163'6-197'6 117'4 and 149'5 199'9 and 250'2 136'5-301'2 214'5-232'2 55'5	1'317 - 4'73 0'1388- 1'623 0'099 - 3'705 0'0663- 4'294 0'114 - 0'178 0'973 - 7'64 0'148 - 6'456 0'0607-10'193 0'5 - 2'01 0'1173- 0'495 0'112 - 7'97 0'0573- 0'4312 0'0164- 4'02 0'098 - 0'2466 0'0308- 0'1863 0'0911-10'24	2·82 -2·469 2·85 -2·798 2·96 -2·855 2·72 -2·89 2·63 -2·24 2·76 -2·846 2·54 -2·489 1·25 ·1·363 2·24 -1·889 3·09 -2·39 2·62 -2·794 2·44 -2·17 3·01 -2·717 2·28 -2·792 2·525·1·814

formula $C = \frac{0.02 \text{ } T^2}{l}$, the atomic lowering C was calculated to be

3°, a value not very far from that obtained in a number of cases.

A few metals appear exceptional. Ramsay remarked that tin, lead, and bismuth have a tendency to become associated, whilst, from the freezing point measurements, aluminium may have a molecule Al₂. From the known tendency of the molecules of elements to increase in complexity as one passes from left to right in the Periodic Table (see p. 12) such association is possible in the case of these elements but there is little other ground for such a conclusion. As will be shown in the succeeding chapter, some of the abnormal results and deviations from the calculated freezing point depressions can be explained either on the ground of combination in solution, or by the formation and separation of solid solutions. Thus, the decreasing molecular weight values with increase of concentration found for sodium and potassium by Ramsay, Heycock and Neville and Tammann may be explained by formation of compounds.* By other

^{*} See also footnote on p. 38.

methods, it has been found that combination occurs of mercury-with sodium, potassium, and thallium ⁵⁷; of tin with aluminium and copper ⁵⁸; and of sodium with potassium, cadmium, thallium, lead and bismuth. ⁵⁹ Solid solutions are also known to be formed between antimony and tin ⁶⁰ and cadmium and mercury. ⁶¹ When these disturbing factors are taken into account, the general conclusion that metals in metallic solvents have monatomic molecules rests on very strong evidence.

(b) Non-metallic Elements.—Tests of the molecular state of these elements have not been numerous. Several gaseous elements, hydrogen, nitrogen, oxygen, appear to raise the freezing point of some solvents,* like benzene, possibly due to separation with the solid. 62 Iodine has been the subject of repeated investigation on account of its tendency to combine with certain types of solvent. 63 In neutral solvents, such as bromoform, it has a molecular weight near 254, corresponding to I_2 . 64 Only phosphorus, sulphur, and selenium, so far investigated, have molecules greater than diatomic, the extent of the association being illustrated by the following values, in which the concentration C is expressed as grams per 100 grams of solvent.

Phosph in Carbon I		Sulphi in Carbon I		Seleni in Phos	
c.	M.W.	C.	M.W.	C.	M.W.
1.281	129	1.245	247	0.503	624
3.723	134	3.123	255	0.340	599
7.44	142	6.14	270	0.413	586
10.84	150	9.04	279	0.480	614
18.86	170	15.13	297	$Se_8 =$	634
$P_{4} =$	124	21.00	314		
_	·	$S_8 =$	256		

Quite a number of investigators have busied themselves with the problem whether the various forms of solid sulphur exhibit different molecular weights in solution, but without finding any such difference. As already stated, two types of molecule, S_8 and S_6 , apparently exist in the vapour near the boiling point of the element. A study of the graphs on p. 18 for the vapour suggests that the amount of S_6 at such temperatures as those at which solution measurements are made will be very small, even in dilute solution (see also chap. VIII.).

^{*}Solvents used in freezing point determinations are sometimes dried by passing through them a current of dry air, which usually has the effect of raising the freezing point.¹¹⁹ In view of Garelli and Falciola's results, care ought to be taken in applying the method and, in any case, on account of the considerable solubility of gases in many organic liquids, the gas should finally be expelled from the solvent.

Once again, the only elements which are known to be associated, are those which tenant Groups V and VI of the Periodic Table, and this fact illustrates the close analogy between gases and solutions.

The Molecular Complexity of Compounds.

The study of the molecular state of compounds has brought to light many substances which exhibit molecular association. A table both of these compounds and of elementary substances, has been compiled as an appendix illustrating not only the variation of the molecular weight with the nature of the substance but also the effect of the solvent in very many cases. Many of the points raised in this and the succeeding chapter are illustrated in this table, making it unnecessary to introduce very complete illustrations in the text, so that it remains only to deal with the results in a general way.

It was found early in the study of the subject that associated substances admitted of ready classification. Associated organic substances, it was found, owe their condition to the presence of certain electro-negative groups. Such groups are the OH, COOH, CONH, NOH, CN, SH, CSNH, CSOH, NH₂, NO, CHO, CO, and CS, conferring the power of association on alcohols, phenols, carboxylic acids, amides and anilides, oximes, nitroso compounds, aldehydes and amides, primary amines, nitroso compounds, aldehydes and ketones. The Appendix contains numerous illustrations of such substances. Solution measurements, therefore, have very greatly enlarged our knowledge of molecular association, as, with the exception of acetic acid, no organic substances are definitely known to be associated in the state of vapour.*

Among the first inorganic compounds to be recognized as associated were the oxides nitrogen peroxide, 68 phosphorous oxide in benzene 69 and arsenious oxide in nitrobenzene. 70 More recently hydrogen chloride and nitric acid and a large number of salts have been added to the list. In the author's view, electrolytes as a class are associated substances, and some evidence of this will be forthcoming in a later chapter.

Dealing now with the substances in three divisions we have :—

^{*}Schulze 67a concludes from his experiments that carbon disulphide and nitrobenzene are slightly associated in the vapour state,

A. ORGANIC COMPOUNDS, EXCLUDING SALTS.

The examples are numerous. It does not follow that one of the electro-negative groups already mentioned confers the power of forming aggregation on the substance without relation to the rest of the molecule. The majority of associated substances contain an electro-negative group united to some hydrocarbon radicle, and as hydrocarbons as a class have no tendency to association, we may expect hydrocarbon radicles to have a normalizing tendency, its extent depending on their mass. Hence, it follows that as a series of homologous substances is ascended, the degree of association diminishes and is ultimately reduced to zero. Thus, methyl and ethyl alcohols, formic and acetic acids, formamide and acetamide are the most strongly associated of their class.⁷¹

Of the groups causing association, some exert a powerful, others only a weak influence. Thus, from the Appendix it will be found that the class of compounds in which the highest degree of aggregation is reached is the alcohols; then follow amides, carboxylic acids, oximes, and anilides. To the oxygen atom must be ascribed considerable power of producing association, for when replaced by sulphur, the analogous compounds containing the groups SH, CSNH, and CS are but slightly associated. Then, again, the groups CHO and CO have also only a weak influence; for whereas acetone is slightly associated in benzene solution, acetylacetone and most other ketones are normal. A number of general rules may be formulated in connexion with the association of organic compounds, and a short statement of the important ones will assist the reader in a survey of the subject.

- I. The tendency to association in organic compounds is connected with the presence of certain electronegative groups, OH, CONH, COOH, NOH, CN, SH, CSNH, CSOH, $\rm NH_2$, NO, CHO, CO, CS, their influence decreasing in the order given.
- 2. The degree of association depends also on the mass of the molecule, diminishing as the formula weight increases. This applies both to aliphatic and aromatic compounds.
- 3. With aromatic substances, the ortho-substituted compound is least and the para-compound most associated. 75
 - 4. The degree of association depends also on the presence of

other groupings in the molecule. Thus, the groups acting as substituents in a molecule, as in the phenols, exert a normalizing tendency as follows:—

$$CH_3 > Cl > Br > I > NO_2 > CO_2R.^{75}$$

- 5. The degree of association depends on the structure of the molecule. Thus, primary alcohols are most and tertiary alcohols least associated.⁷⁶
- 6. The degree of association depends on the solvent (chap. IV.).

B. SALTS.

By far the most systematic comparison of salts has been carried out with those derived from organic ammonium, sulphonium and oxonium bases, as these dissolve in a wide range of solvents much more readily than metallic salts. Strong association has been found with all true salts, the degree of association being, on the whole, considerably greater than with non-electrolytes. Thus, in chloroform, tetraethylammonium bromide may have, according to the concentration, a degree of association greater than eight, and triethylsulphonium iodide of twelve. The actual degree of association depends on a number of factors, 77

I. On the base.—As a rule, the degree of association of similarly constituted salts decreases as the mass of the base increases. Thus, the association decreases in the order monoalkyl-, dialkyl-, trialkylammonium chlorides; dimethyl-, diethyl-, and dipropylammonium chlorides; tetraethyl- and tetrapropylammonium chlorides. But mere mass is not the controlling factor, and structure must play a considerable part; for, instead of a decrease in the degree of association being found on passing from the trialkyl- to the tetraalkylammonium salts, an increase occurs; likewise, the quaternary compound, quinoline ethiodide, is more associated than quinoline hydriodide.

So far as the alkylammonium and alkylsulphonium bases are comparable, the salts of nitrogen bases are the more strongly associated.

2. On the acid.—Of the halogen salts, chlorides appear least and iodides most strongly associated, so that the degree of association does not fall, here, but rises with increase in mass. Indeed, with periodides, 78 the extent of complexity appears to be

still greater than with iodides. Nitrates 77 are likewise strongly associated, in chloroform slightly more than iodides.*

3. On the solvent, the concentration and the temperature (see below).

For metallic salts, very similar rules hold,⁷⁸ the degree of association falling with rise of atomic weight for compounds of allied elements, whilst the chloride is also least and the iodide most associated of the haloid salts, as in the lithium compounds.

The tendency to molecular association shown by inorganic substances appears to be connected with the relative positions of the component elements in the Periodic Table. Nitrogen peroxide, phosphorous oxide and arsenious oxide are compounds of oxygen and elements in Group V. All of them are non-electrolytes, though, as anhydrides, they can readily produce electrolytes by the addition of water. The electrolytes are strongly associated, as hydrogen chloride, which is strongly associated in benzene and nitrobenzene, 119 and so also are salts. The essential difference in the behaviour of electrolytes and non-electrolytes is illustrated in the compounds of the halogens with Group V elements. phosphorus and arsenic tri-chlorides are non-associated, but antimony trichloride exhibits quite distinct association. evidence indicates that as we pass from left to right in the Periodic Table, the power of the haloid compounds of the elements to form molecular aggregates diminishes.⁷⁹

Solid Solutions.

In the region of solid solutions the most available method is the application of the Partition Law. So far as the discovery of association, or any abnormal molecular complexity is concerned, the results are as yet very meagre. One of the most interesting cases concerns the absorption of gases in solid materials. When hydrogen diffuses through palladium 80 or through iron, 81 or dissolves in palladium, 82 or nickel, 83 it does so to an extent which is proportional to the square root of the pressure. There are two possible ways of explaining such a result, either that Henry's Law in regard to the solubility of gases (namely, that $\frac{C_1}{C_2}$ shall be constant, where C_1 measures the concentration (pressure) of

^{*} Fluorides, chlorates, bromates, iodates, and sulphates have also been found, from measurements in the author's laboratory, to be associated strongly.

the gas outside the absorbing medium and C2 that within the medium), does not apply to such cases of absorption; or, that the gas must diffuse in the atomic state, when $\frac{\sqrt{C}}{C}$ should be constant. One difficulty in the way of the second explanation lies in the fact that the absorption of carbon dioxide in charcoal 84 is also expressed by the relation $\frac{x}{\sqrt[3]{P}}$, from which it might also be concluded that the carbon dioxide molecule was dissociated into three parts during the process of absorption. explanation for carbon dioxide is not thinkable. It is now generally recognized that for most gases, Henry's Law does not apply to their absorption by solids, but that in at least one case, hydrogen in charcoal, there are two processes, the first a rapid absorption to which Henry's Law does not apply, the second a very slow process of true solution. According to McBain, 85 the second process of true solution occurs at a rate proportional to the square root of the pressure, and the only conclusion to be drawn here is that the hydrogen suffers a dissociation, diffusing inwards in the atomic condition. Not only is this conclusion drawn for solid solution, but also for solutions of hydrogen in molten copper and of oxygen in silver and in gold-silver alloys.86 Sieverts and Hagenacker suggest that such dissociation into the atomic condition may be occasioned by the monatomic molecules of the metal in some way. Such an explanation cannot be applied, however, to solution in charcoal.

Solutions of solids in solids so far investigated, include no case in which the degree of association has been measured. If a solute is associated in a given solvent and separates with the solid solvent in a manner governed by Henry's Law, then the conclusion may be drawn that it is associated to the same extent both in the liquid and solid solutions. Probable examples are furnished by antimony and bismuth bromides in aluminium bromide, but the partition coefficient between liquid and solid has not been satisfactorily determined in these cases.

Colloidal Solutions.

Colloids furnish another and very interesting type of association. They are substances which possess a very big molecular weight, as all results indicate. Thus, Pfeffer found that a solution of gum arabic in water, at 15°, exerted but a small osmotic pressure, the values of the molecular weight calculated from the osmotic pressure ranging from 2570 to 4110,88 but obviously without any great degree of accuracy.

Freezing point measurements lead to very similar results. In water, quite a number of cases have been tested, of which the following table includes a selection:—

Concentration.	M.W.
14.2-44.2	14,000
10.0	6500
?	25,000
14.0	1612
8.76 - 22.5	1585-1745
1.16 - 5.60	5452-4888
0.523 - 1.37	410-1073
1.2 - 100	679- 995
-	800-1600
	14'5-44'5 10'0 ? 14'0 8'76 - 22'5 1'16 - 2'60 0'523 - 1'37

The objection to freezing point measurements is the possible influence of small quantities of salts or other impurities which, on account of the adsorptive power of colloids, it is difficult to remove; and the question has been raised whether colloidal substances of themselves can exert an osmotic pressure. Despite certain evidence that no osmotic pressure is exerted, ⁹⁴ it can be safely concluded from the work of Linebarger, Lillie, Moore and Roaf, ⁹⁵ and others that a definite though small osmotic pressure is set up, corresponding to a high molecular weight. ⁹⁶

Although the investigation of the conditions of the formation of colloidal solutions is far from complete, there are certain resemblances between ordinary association and colloidal solutions which may be indicated. In the first place, the gas laws and laws of distribution which apply to substances in ordinary solution, whether associated or not, have more recently been shown to be equally valid for colloidal substances.⁹⁷ We may in fact regard the formation of colloidal solutions as a limiting case of association, and it is quite possible that the upper limit of the degree of association of, for example, salts, would be found in a colloidal solution. Already, from measurements made in chloroform, the molecular weight values for some salts run up to numbers which one is accustomed to connect with colloidal substances as, for example, tetraethylammonium bromide which reaches (not as a limit) a value exceeding 1800, or of triethylsulphonium iodide exceeding 2500.98 Then again, the influence of the solvent, in a very general way, is the same in both cases. It has been pointed out elsewhere, that association of a substance occurs as a rule in solvents in which the substance is but little soluble. This applies also to colloidal solutions, for colloids are produced in solvents in which true solution is not formed. Thus, ordinary soluble salts are looked on as crystalloids in regard to aqueous solution. At any rate they do not form colloidal solutions in water* but can do so in organic liquids in which they are insoluble in the ordinary sense. Arsenious and antimonious sulphides which are insoluble in water, are associated in the sense of forming colloids, in this medium. What the effect of a series of solvents is on the size of colloid particles cannot be said. Organic substances can form colloidal solutions in media other than water, as, for example, caoutchouc in benzene 100 and casein and starch in formamide. 101

The effect of temperature increase is also to reduce the size of colloidal particles.¹⁰² One further factor, the influence of added electrolytes is, on the other hand, to increase the size of the colloidal particle, or to diminish it according to the character of the ions added, before precipitation occurs, as has been proved by Biltz and Vegesack, and Mayer, Schaefer and Terroine.¹⁰³ Little is known of any similar phenomenon connected with ordinary association, but the recently demonstrated effect of one salt increasing the degree of association of a second in solution,¹⁰⁴ may, at any rate in the early stage of the action, have something in common with the effect of electrolytes on colloids.

The Influence of Temperature on Molecular Weight.

The range of temperature over which it is possible to study the molecular weight of a substance in solution is in most cases much more limited than with a gas. Arguing from the analogy between the two states, it would be expected that rise of temperature should bring about a reduction in the degree of association of a substance, and so far as tests have been made this actually is the case. With a solution, however, it must be clearly borne in mind that other factors have to be considered. Thus, a very important influence is exerted by the dielectric constant of the solvent, and the value of this property decreases with rise of temperature,

^{*} Unless they are protected by a coating of a natural colloid as in the case of silver chloride. 99

tending, therefore, to maintain the associated condition. ¹⁰⁵ To ascertain the true effect of temperature, therefore, a knowledge of the temperature variation of other factors is obviously essential.

There are several ways in which the influence of temperature may be studied. The first is to make determinations of the molecular weight by freezing point and boiling point methods for a given solvent. Thus we have the following values derived in this way, the concentration being, for benzoic acid in benzene, 25 milligram molecules per 100 grams of solvent; for tetraethylammonium bromide, 25 milligram molecules per 100 c.c. of solvent.

BENZOIC ACID IN	BENZENE,106
to	M.W.
5°5°	233
8o°	193

Tetraethylammonium Bromide in Acetic Acid. 107

16.6° 527 118° 337

A more thorough test was that made by Innes 108 on the degree of association of several organic substances, namely, benzoic and o-bromobenzoic acids, β -benzilmonoxime and dimethyltartrate in the solvent benzene, which was made to boil at a series of temperatures by varying the external pressure. A series of measurements was first carried out with normal substances in order to deduce the molecular elevation of boiling point at each temperature. Fig. 4 records graphically the results with benzoic acid and it will be seen that with rise of temperature there is a steady fall in the molecular complexity for any given concentration. The curve for 93° lies above that at 80° and in its upper portion cuts the curve for 73°, but the known volatility of benzoic acid below its melting point may account for this. The other substances tested gave results of similar character.

As already mentioned, other factors, the variation of the dielectric constant must be remembered. In the case of benzene this variation is, however, very small.*

The heat of association or dissociation must be a factor determining the stability of the molecular complexes. Its actual value could be determined, as with gases, if the precise effect of temperature on the degree of association were known. 108

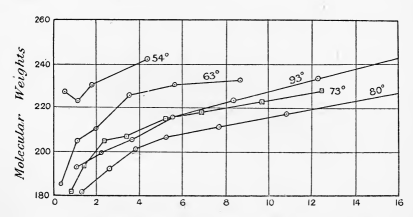
^{*} As yet unpublished result.

The Effect of Concentration.

As already seen in chapter II. cases of molecular association follow, in the simplest case, the equation :—

$$A_n \geq nA$$

and must be influenced, accordingly, by the pressure or molecular concentration in such a way that increase of concentration brings about an increase in the degree of association. Where experiment proves that the reverse occurs, namely, decrease with increasing concentration, it can with considerable assurance be concluded that chemical combination occurs between solvent and solute.

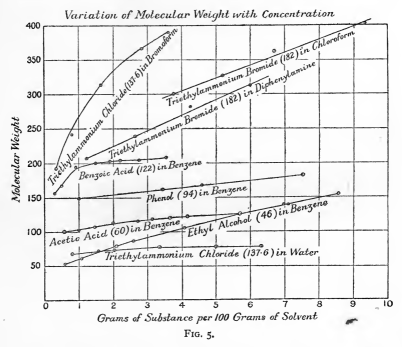


Hundredth-gram Molecules per 100 grams of Solvent. Fig. 4.

Fig. 5 indicates for some typical cases the manner in which the molecular weight varies with concentration. In general, a particular, form of curve is representative of a class. Thus, Auwers ¹⁰⁹ found that the alcohols have nearly straight line curves, rising rather steeply, and so have the phenols, amides, and anilides. Salts also generally follow this type. ¹¹⁰ Benzoic and similar aromatic acids are strongly associated even in dilute solution, and after rising steeply for a short distance, the curve rapidly changes in direction and its slope becomes quite small.

The slope of the curve will obviously depend on the limiting value of n in the complex A_n . Auwers was led to divide associated substances into two classes on the basis of this difference in the type of curve, the first class comprising the aromatic acids, in which there appeared from his measurements to be an upper

limit of n=2, the second class containing substances in which the molecular weight increases proportionately to the concentration. Most substances belong to this second class, alcohols, phenols, aliphatic acids and salts. Views have been expressed that some salts have an upper limit to the degree of association. Thus, Hantzsch and Hofmann ¹¹¹ believed that tetraalkylammonium salts in chloroform behave very simply, the molecular state being nearly independent of the concentration and having a degree of association of five as an upper limit; whilst Wedekind



and Paschke ¹¹² regarded salts as having an association factor of 2. Both deductions were made from insufficient data, for, to quote but one example, tetraethylammonium bromide in a 9 per cent solution in chloroform has a degree of association of 9. Certain salts appear unaffected by alteration of concentration, but the molecular weight found in such cases is often nowhere near a multiple of the simple molecular weight.¹¹³

What the upper limit of complexity may be in the case of substances which at moderate concentrations have straight line curves connecting molecular weight and concentration is rather an interesting speculation. The ordinary methods available for dilute solutions could not be applied.

When we come to deal with the lower limit of association we are on surer ground. Continuous reduction of pressure on a gas tends to reduce the molecular state to the simplest possible, and in like manner, decrease of concentration should bring about a similar state in solution. As will be seen from Fig. 5, there are one or two cases in which expectation is fulfilled, for example, ethyl alcohol in benzene. On the other hand, the production backwards of several of the curves would not lead to intersection of the y axis at the normal value of the molecular weight. Probably, however, the direction of the curve for very dilute solutions alters as the concentration increases.

The quantitative study of the relation between concentration and degree of association has, however, made but little progress. On the basis of the Law of Mass Action, assuming a direct change into single molecules on dissociation, the state of equilibrium $A_n \gtrsim nA$ must be expressible by the equation

$$K = \frac{C_A^n}{C_{An}}$$

where C_A and C_{A_n} are the respective concentrations of single and complex molecules. Nitrogen peroxide in chloroform solution, affords the only satisfactory example of an associated substance which dissociates in solution according to the requirements of the Law of Mass Action.^{113a}

The resemblance between the curves representing the dissociation of benzoic acid in benzene and of weak electrolytes in water led Auwers 114 to apply the equation to substances of the first class, namely, aromatic acids, for here, presumably, the limits are simple, varying from n = 1 to n = 2. The equation then becomes

$$c = K \frac{I - y}{y^2}$$

in which c represents the concentration, y the degree of dissociation of the double molecule and K the equilibrium constant.

If, therefore, in a solution of unit molecular concentration, calculated on the basis of the single molecule, e represents the number of single molecules, d the number of double ones, then

$$e = I - 2d$$
 or $d = \frac{I - e}{2}$.

But the lowering of freezing point and the number of molecules are so connected that

$$\frac{m}{M} = e + d$$

from which

$$d = \frac{M - m}{M}$$
, and $e = \frac{2m - M}{M}$.

Further,

$$y = e = I - 2d = \frac{2m - M}{M}.$$

Substituting these values in the simple mass action formula, we get,

$$K = c \frac{(2m - M)^2}{2M (M - m)}.$$

This formula was applied to various cases of association but only found to be in any way satisfactory in very few cases, as with benzoic acid in benzene and o-bromobenzoic acid in naphthalene, for which the following values were calculated:—

Benzoi	c Acid (m	= 122).	o-Bromober	zoic Acid	(m = 201)
C.	M.	K.	c.	M.	K.
0.568	219	0.0039	0.50	269	0.14
0.262	223	°0056	0.48	293	0.12
1.444	228	•0076	1.26	311	0.19
2.603	232	*0073	3.02	332	0.12
4.725	236	·0056	4.21	340	0.18

For most acids as well as oximes and members of the second class, the values of K decreased throughout with increasing concentration, and even the aromatic acids only gave satisfactory results when the concentration did not exceed 4 centigram molecules per 100 grams of solvent. The phenomenon of association in solution is evidently not simple. Possibly some of the factors which cause strong electrolytes in aqueous solution to disobey the Law of Mass Action also operate in complicating the process of association. It is quite possible that the assumption of an upper limit of 2 for the degree of association is not correct and that more than two types of molecule may be present, in which case each dissociation will be marked by a distinct equilibrium constant. Auwers himself suggested the cause of the disturbance to be some reaction between solvent and solute, an idea which he proceeded to test by investigations on the influence of the chemical constitution of the solvent, an influence which he found quite as powerful as the nature of the solute itself. The whole subject of this influence will be dealt with in a general way in the succeeding chapter.

CHAPTER IV.

THE MOLECULAR COMPLEXITY OF DISSOLVED SUBSTANCES.

The Influence of the Solvent.

So far in our study of molecular association, the known analogies which exist between the gaseous state and solution have been of considerable assistance; but now new factors must be taken into account. For although it is interesting to liken a gas to a solution in the ether (assuming the existence of the ether), little is known of such a medium beyond its power of conveying energy. In the measurement of gas densities no second substance is present, except in the methods of Meyer and of Lumsden; but here the only effect which the filling gas exerts is due to its action as a diluent, bringing about somewhat increased dissociation, where dissociation can occur at all, to an extent depending on the extent of admixture with the vapour. In any case, the filling gas is so chosen that chemical action is rigidly excluded.

In a liquid solution, however, the solvent plays an important part. Its action may be wholly physical, or partly also chemical. The effect of chemical action will be considered first.

I. Combination between Solvent and Solute.

At first sight, there are two possible ways in which the observed molecular weight of a substance may be affected by combination in solution. On the one hand it may be argued that the compound formed will make its presence known in an increased molecular weight, the value found being due partly to uncombined solute and partly to the compound; on the other hand, the action may be regarded as merely reducing the amount of free solvent, thereby concentrating the solution and leading to a lower molecular weight, since the osmotic pressure, depression of freezing-point, or vapour pressure, will be thereby increased. Ramsay, 115 in his study of metallic solutions, took the former

view; the result of experiment, however, may be said to prove the correctness of the second.

The proof of the reduced value in the molecular weight observed when combination occurs lies in the study of cases in which combination is known to occur.* It has recently been pointed 116 out that, although the general effect of combination is always to make the observed molecular weight lower than it would be in the absence of such combination, there are five ways in which the molecular weight may vary with concentration, supposing molecular association and combination both to exist.

I. The molecular weight decreases with increasing concentration.—Benzoic acid in p-toluidine, ¹¹⁷ for example, where combination occurs, illustrates this case. Another example, of a different type, is that of lithium chloride in ethyl alcohol. ¹¹⁸ These two substances combine to form LiCl, 4C₂H₆O. The variation of the molecular weight is as below, C indicating the number of grams of the salt dissolved in 100 grams of the alcohol:—

C.	M.W.	
2.244	34.0	
2.655	33.0	LiCl = 42.5
3.406	30.1	
4°075	29.2	

Hydrogen chloride, both in benzene and in nitrobenzene, is strongly associated, but as the concentration increases, the apparent degree of complexity falls off, a fact only to be explained by the occurrence of chemical combination.¹¹⁹

- 2. The molecular weight increases with increasing concentration.—Here, despite combination, the rate of increase of complexity is greater than the reducing effect due to combination. Several examples occur in the determinations of Werner, such as in the case of cadmium bromide in piperidine. 120
- 3. The molecular weight increases to a maximum and then falls.—Such a phenomenon will be observed in cases where there is a rapid increase in the degree of association in the early stages
- * Neglect to take into account the heat of dilution introduces the same kind of an error as is now discussed under the head of chemical combination. Thus, when the heat of dilution is taken into account, the molecular weights of sodium and potassium in mercury (p. 23) have normal values. It is probable, however, that in most cases where it is necessary to correct the osmotic pressure (freezing point, etc.), for heat of dilution, that chemical action between solute and solvent occurs; so that a chemical explanation can, in fact, be offered for an abnormal physical property.

For a discussion of the effect of heat of dilution see Findlay's "Monograph on Osmotic Pressure" in this series.

(that is, while the solution is still dilute) since the effect of combination only becomes pronounced in concentrated solution. Quite a number of examples may be quoted; indeed, this type seems by far the commonest. Lithium iodide in amyl alcohol, from which, at lower temperatures LiI, $4C_5H_{12}O$ can be isolated, and potassium iodide in liquid sulphur dioxide, where combination to form KI. $4SO_2$ occurs, may serve as examples. C represents grams of solute per 100 grams of solvent.

Lithium Iodide in	Amyl Alcohol.118	Potassium Iod	lide in Sulphur Dioxide.121
C.	M.W.	C.	M.W.
1.777	154.4	0.371	282
4.122	159.2	o·735	331
8.053	151.2	1.13	352
11.97	151.2	1.28	288
		2.05	289
		2.46	222

Other examples are furnished by lithium bromide and iodide in butyl alcohol and lithium bromide in amyl alcohol, where compounds of the type LiBr, 4ROH are formed; ¹¹⁸ by potassium iodide in iodine, ¹²² and probably a variety of salts in p-toluidine. ¹⁰⁵

The possibility of the two other cases follows from the three already given, so that it is quite possible to have (4) the molecular weight remaining constant over a certain range of concentration, the increase of association being counterbalanced by the effect of combination, and (5) the molecular weight falling to a minimum value.

The fact to be borne in mind is that combination in solution has the effect of reducing the molecular weight. Further evidence, beyond that given, is that where combination does not take place, the molecular weight of an associated substance increases steadily with the concentration, as with lithium nitrate in ethyl, *iso*-butyl, and *iso*-amyl alcohols, where no compounds are formed.¹¹⁸

II. The Physical Influence of the Solvent.

A. Separation of the Solute with the Solvent.—This is a factor which is apt to cause confusion, when not understood, and to lead to the idea either that a non-associated substance is associated or that the degree of association is greater than is actually the case. For if the solute separates in any manner with the solvent, the solution remaining is more dilute than if no separation occurred. In consequence, the osmotic pressure, as measured by

freezing point or vapour pressure determinations, is reduced and the apparent molecular weight is high. Such separation can occur either by the formation of solid solutions or by the volatility of the solute in a boiling point or vapour pressure determination.

An interesting example is furnished by iodine in benzene. On other grounds there is no evidence that this element is associated (cp. p. 24), but cryoscopic determinations in benzene lead to molecular weights between 350 and 360, whereas $I_2=254$. The explanation was forthcoming when Beckmann and Stock ¹²³ found that iodine forms a solid solution in benzene.

If the extent to which a substance separates with the solvent is determined this amount can be allowed for and the true amount in solution deduced. Supposing that a is the fraction of the substance in the solid solution or in the mixed vapours, then the concentration of the liquid solution will be reduced to C(I-a), where C was the original concentration. Therefore the usual equation for the calculation of molecular weight becomes:—

 $M = \frac{C (I - a) K}{\cdot \Delta^o W}$, where K is the elevation or depression constant, Δ^o the observed rise of boiling point or fall of freezing point and W the weight of solvent. The quantity a corresponds to the partition co-efficient, and can be determined by a method such as that of Bijlert, 124 for solid solutions. The determination of a in the vapour state is less easy. If the solute is an acid, however, the vapour arising from the solution can be distilled over and analysed. 125

As an indication of the extent of error involved by the separation of solute with the solvent, the behaviour of thiophene in benzene may be quoted. 126

C.	M.W.
0.21	133
1.13	133
2.16	133
3°25	134

When correction was made for the amount of thiophene freezing out with the benzene, the mean molecular weight observed became 78·I as compared with the calculated value of 83·8. When a non-associated solute separates with the solvent, therefore, the observed molecular weight is higher than the normal, but is independent of the concentration. The degree of com-

plexity of an associated substance is, as already indicated, dependent on the concentration.

How many of the cases of actual association are subject to an error due to separation with the solvent is quite unknown, as it is seldom that the nature of the solid separating out is tested. Acetic and benzoic acids and a-benzaldoxime in benzene certainly are not. Benzoic and a number of aromatic acids do separate with water vapour at the boiling-point of the solvent, and antimony and bismuth bromides in aluminium bromide are further examples of associated substances which separate with the solvent, in this case on freezing. 87

B. Dielectric Constant of the Solvent and Other Physical Actions.—The power of forming complex molecules in solution appears to depend in a very great measure on the solvent apart from any chemical action. Raoult and the early workers in the same field found that the association of organic substances containing the hydroxyl group was exhibited when the solvent was a hydrocarbon or a halogen or nitro-derivative of a hydrocarbon, and all subsequent work substantially proves this regularity.

The first systematic researches on the influence of the solvent were made by Auwers who studied the effect of the chemical constitution both of the solute and the solvent on the degree of association, finally coming to the conclusion that the influence of substituent groups could be set out in a regular manner, in the order

OH, COOH, NH₂, CHO, CN, COOR, NO₂, I, Br, Cl.

When these groups are present in the molecule of a solute, the most abnormal result is obtained when the OH group is present, and least with Cl; when present in the solvent, the OH group has the greatest normalizing effect (that is, is most effective in reducing the molecular weight of a solute to the normal) and Cl least. This rule was put forward as expressing in a general manner the results obtained without being claimed as applying universally.⁷⁵ Certain possible physical influences were tested, such as the melting-point and the molecular freezing-point depression of a given solvent, but were found to be without effect.

It is quite possible, however, to refer the molecular complexity more or less successfully, according as the solute is an electrolyte or a non-electrolyte, to the dielectric character of the

solvent. Already, in 1893, J. J. Thomson 127 had put forward the hypothesis that "if the forces which hold the molecule together are electrical in their origin, . . . these forces will be very much diminished when the molecule . . . is surrounded by a substance possessing a very large specific inductive capacity". reference, although in a paper on the conduction of electricity through gases, suggested a connexion between dissociation in aqueous solution and the specific inductive capacity of the solvent. Shortly afterwards Nernst, 128 again on the assumption that chemical forces are electrical in their origin, added the statement in regard to association "Wir werden erwarten können, dass Lösungsmittel um so stärker Doppelmoleküle zu spalten vermögen, d. h. eine um so grössere dissocierende Kraft besitzen, je grösser ihre D. E.* ist. Dies bestätigt sich volkommen". Nernst referred to rather different substances from Thomson, for the association of salts was not then known. The reference, in particular, was to organic acids, which are associated in benzene, a solvent of low dielectric constant. The combined statement of Thomson and Nernst is known as the Nernst-Thomson hypothesis, and until the last few years had been confined in its application to the ionization of electrolytes.

It is not difficult to show that Auwers' results can be referred fairly successfully to the dielectric constant of the solvent. As already stated the normalizing effect of different groups contained in a solvent decreases in the order

OH, COOH, NH₂, CHO, CN (COOR), NO₂, I, Br, Cl. Now the dielectric constant of a solvent is likewise affected, in general, by the nature of the groups present. According to Walden, decrease in the dielectric constant occurs as substitution proceeds in the order from left to right,

OH, NO₂ (CO), CHO, COOH, CN, NH₂, F, Cl, Br, I. The order is not quite the same in the two cases, the NO₂, COOH, and NH₂ groups being in different positions; but as the effect of the halogens on the dielectric constant may also be in the reverse order to that given, there is a similarity between the two arrangements.

The effect of the dielectric constant can be better illustrated by reference to the actual numbers obtained. Weak electrolytes, benzoic acid, and acetic acid, and a non-electrolyte, formanilide,

^{*} The symbols D. E. refer to the dielectric constant.

will be first considered. In the following table the original results, drawn from a large number of sources, have been collected and recalculated to a common concentration, namely, a solution containing 30 milligram molecules of the solute (assuming the simple formula for the calculation) per 100 c.c. of the solvent. M_{30} gives the molecular weight at this concentration. K_t represents the value of the dielectric constant at the temperature t^9 , as near as possible to the temperature of the determination.

		M ₃₀ .				
Solvent.	Kt.	Benzoic Acid (122).	Acetic Acid (60).	Formanilide (121).		
Water	66 ₁₀₀ °	* (240) 125	58 138	157 143		
Ethylene cyanide	61.500	180 130	_	-37		
Nitrobenzene	40.060	280 ¹¹⁹	(associated) 139			
m-Dinitrobenzene	20.6000	167 75				
p-Nitrotoluene	18.4500	_	_	157 75		
Ethyl alcohol	18.0 ⁴⁸ 0	125 131	_	I 17 143		
Acetone	17.0260	110 131		120 143		
Sulphur dioxide	14.0500	222 132	94.6 132			
Phenol	9°748°	133 133	_	-		
Acetic acid	6.3180	120 134	_			
p-Toluidine	5°4 ₄₃ °	142 117	6.710	128.5 75		
p-Bromotoluene	5'υ ₇₂₈ °	_	106 140			
Ethylene dibromide	4.86180		118 139			
p-Dibromobenzene Bromoform	4°5788°	230 75	0.141	_		
	4'37.5°	-66.75	121.8 141			
p-Methyl toluate Chloroform	4.28330	166 75		149 75		
Ether	4'1361.2	214 ¹³¹ 126 ¹³¹	_	160 143		
Naphthalene	4'12350	212 136		164 143		
Nitrogen peroxide	3.5800	210 135	(120) 142	_		
Carbon disulphide	2.2	243 131	(120)	_		
Benzene	2.3 _{5.5} °	236 ¹³⁴	117 134	213 143		
p-Xylene	2.1218.5	230	11/	244 75		
Cyclohexane	2.02	>230 137	135 137	444		

The view of Nernst, based on a very limited number of results, and of Meldrum and the writer, 144 from more extensive measurements on amides and anilides, was that the dielectric constant of the solvent was probably the controlling factor on association. This conclusion, in the case of the substances selected as examples in the above table, requires some modification, and further investigation. It is true that in solvents of low dielectric constant, association almost always

^{*} This is the value at the concentration 50 milligram molecules per 100 c.c. of solvent.

occurs. Ether as a solvent for benzoic acid is obviously an exception, although it falls into line when formanilide is the solute, and the low value with p-toluidine can, in the case of benzoic acid, be referred to chemical action (vide p. 38). But, although association is almost general in solvents of low dielectric constant, it is far from absent when the dielectric constant is high. Thus, pronounced association occurs with benzoic acid in water, ethylene cyanide, nitrobenzene and m-dinitrobenzene, whilst formanilide is associated in water and p-nitrotoluene. It must be conceded that although the dielectric constant undoubtedly plays a distinct part, it is not the controlling factor with all organic associated substances. The influence of the chemical nature of the solute is evident in the three examples chosen, and it appears that there is a distinct difference between aliphatic and aromatic compounds. Acetic acid, so far as has been tested, is much more influenced by the dielectric constant than benzoic acid and formanilide, which contain the benzene nucleus. The examples quoted in the Appendix add further evidence of this fact.

The existence of molecular complexes in aqueous solution is hardly to be looked for on the usual view of the dissociating effect of water, but such is actually the case. A number of organic acids and phenols, 145 amides and anilides 146 appear to be strongly associated in water as well as in the so-called associating solvents, a term whose application becomes difficult in view of the effect of water as a solvent. The occurrence of molecular association in water is of interest and importance from other points of view, however, for in practically all cases, the more strongly associated the substance in benzene and solvents of low dielectric constant, the less the substance is associated in water, and vice versa. 146 A few examples will make this clear, the association factors referred to being deduced for a common concentration of 50 milligram molecules per 100 c.c. of solvent.

	Associa	Association Factor.		
Substance.	Water.	Benzene.		
Propionamide	1.14	2'03		
n-Butyramide	1.13	1.08		
iso-Butyramide	1.18	1.94		
Valeramide	1.22	1.82		
iso-Butylacetamid	e 1.20	1.49		
Methylacetanilide		1'04		

			Associa	tion Factor.
Su	bstance.		Water.	Benzene.
	oic acid		1.08	1.97
o-Hy	iroxybenz	oic acid	1.40	
m-	,,	,,	1.53	
p-	,,	,,	1.14	
Phen			1.13	1.89
p-Nit	rophenol		1.46	1.38

The hydroxy-acids quoted are not sufficiently soluble in benzene to be tested, but from tests in naphthalene it appears that the usual rule of the para- compound being most associated and the ortho- least is adhered to. The inversion of the order for benzene and water is very clear. Once more the strong effect of the benzene nucleus in producing association in spite of the solvent is brought out.

On the basis of the regularity demonstrated, the suggestion has been made that possibly the association in water is only apparent, and that in reality it is due to the dissociation of the water molecules, which are known to be complex ^{145, 146}. A number of other observers, Robertson, ¹⁴⁷ Armstrong, ¹⁴⁸ and Sutherland ¹⁴⁹ have made similar suggestions based on other types of measurement. If the complex water molecules are dissociated, then the concentration of the solution in respect to the solute is diminished and the molecular weight from the observed osmotic pressure becomes greater. Such a mode of explanation is not easy to test, and it must be pointed out that association of benzoic and acetic acids occurs in nitrobenzene, a substance whose molecules are simple, according to current conceptions.

On the other hand, if the association in water is real and not apparent, it may be of the same type as that which is exhibited by colloidal substances; and this at any rate can be said for the view, namely, that the substances on the whole least soluble in water are the ones most associated, just as colloids do not dissolve to form a true aqueous solution (cp. p. 31). But supposing this to be the case, that is, suppose the association in water to be just as real as in any other solvent, then a formidable difficulty of another kind arises. Nernst 150 based his theory of the Partition Law both on examples of normal substances and on the instance of benzoic acid divided between water and benzene. Now, on the assumption that benzoic acid is normal in water and has approximately a double molecule in benzoic acid, Nernst

found that the requirements of the law that $\frac{C_w}{\sqrt{C_R}} = k$ (where C_w

is the concentration in water and C_B in benzene), were fulfilled. From the measurements quoted above on the state of benzoic acid in water, it would appear that in this solvent also the molecules are double, and therefore it would be impossible for $\frac{C_w}{\sqrt{C_B}}$ to be constant. In such a case, the simple ratio $\frac{C_w}{C_B}$ ought to be constant. Experiments made in the writer's laboratory at 40° in order to obtain stronger solutions of benzoic acid, showed the constancy of the ratio $\frac{C_w}{\sqrt{C_B}}$. The problem is, therefore, at present at an interesting stage.

The Effect of Dielectric Constant on Electrolytes as Solutes.

Although the recognition of electrolytes as associated substances is but very recent, the effect of the solvent can be set out more satisfactorily and with greater sureness than in the case of weak electrolytes or non-electrolytes.

The chief difficulty in testing the effect of the solvent on the molecular complexity of salts (for these, together with a few acids, are the only electrolytes yet tested), is their sparing solubility. This difficulty can be overcome, however, by using salts derived from organic ammonium, sulphonium and oxonium bases; for such salts have been found in other respects to behave like the salts of the alkali metals, and so far as metallic salts have been tested, the general regularity shown in the influence of the solvent on organic ammonium salts holds also for metallic salts. With this remark, and reference to the Appendix, the examples chosen for illustration in this chapter will be confined to the salts of organic ammonium bases. 162

The following table sets out for comparison, in a manner like that for non-electrolytes, the degree of complexity of the four salts diethylammonium chloride, tetraethyl ammonium iodide, tetrapropylammonium nitrate and quinoline ethiodide at the same molecular concentration (calculated from the simple formula), namely, 25 milligram molecules per 100 c.c. of the solvent. Instead of quoting the actual molecular weight, the degree of association, or ratio of the observed to the calculated molecular weight is given, and where the number is enclosed in brackets, it

is obtained by extrapolation and consequently subject to possible error.

DEGREE OF ASSOCIATION IN SOLUTION.

CONCENTRATION, 25 MILLIGRAM MOLECULES PER 100 C.C. SOLVENT.

Solvent.	Dielectric Constant.	Et ₂ NH ₂ Cl.	Et ₄ NI.	Pr ₄ N. NO ₃ .	C9H7NEtI.
Formamide	>8420°	0.22	0.58		_
Water	66,000	0.61	0.66	0.66	0.01
Ethylene cyanide	61.5	_	0.63		_
Formic acid	58.5160	0.68	o.68	_	_
Nitrobenzene	40°0 ₆ 0	_	_	(0.92)	_
Nitromethane	39*420°	_	0.68	_	_
Acetonitrile	30810	1.68	0.40	0.63	l —
Propionitrile	22.5980	_	0.77	_	_
Methyl alcohol	21.0660	_	0.80	_	_
Ethyl alcohol	18.0780	1.00	0.99	0.89	1.13
Acetone	17°0 ₅₆ °	. —		1.12	_
Sulphur dioxide	14°O ₂₀ 0	1.41	0.78	_	_
Ethylurethane	13.600	1.27	-	1.00	1.33
Phenol	9.748°	1.24	1.53	1.22	_
Pyridine	9.51160	1.83	(? 2)	_	_
Dimethylethylcarbinol	7.91020	1.54	'	_	
Acetic acid	6.3 ¹⁶ °	1.63	_	2.13	_
iso-Amyl alcohol	5.7132°	1.48	1.87	1.73	2.29
p-Toluidine	5'444°	1.22	1'45	1.21	1.67
Bromoform	4.37.50	3.03	_	_	
Chloroform	4.1361.50	2.23		5.0	(? > 4)
Diphenylamine	3'3 ₅₁ °	1.08	5.0	3.98	

The results in all cases (and the salts chosen for illustration are but types) indicate that the observed molecular weight is very low in solvents of high dielectric constant, rising steadily with decrease of the dielectric constant until, when the latter becomes small, the molecular weight is very high. This transition would certainly be more regular if the determinations were made under precisely the same conditions of temperature, conditions impossible of attainment. Thus, the tests in acetic acid were carried out at 16° while those in amyl alcohol were made at 132°, a difference of temperature which will certainly account for the break in the order given (cp. p. 32). Again, the lower values with p-toluidine are most probably due to combination in solution. On the whole, therefore, it can be said that for salts, the apparent molecular weight is controlled mainly by the dielectric constant of the solvent.

Specific influences, due to the nature of the solute and of the solvent, are not absent. Thus, diethylammonium chloride, and iso-butylammonium chloride, are both distinctly associated in acetonitrile, of high dielectric constant, whilst the same remark

applies to hydrogen chloride in nitrobenzene. ¹⁵³ In the case of the solvent, influences which cannot be ascribed to the dielectric constant or to the employment of different temperatures are not absent, as illustrated by a comparison of chloroform and diphenylamine.

In the light of the preceding results, the abnormally low molecular weight values found for salts in water, results on which the theory of electrolytic dissociation was largely based, no longer stand out as abnormal, but fall into line with those obtained in other solvents. Electrolytic dissociation and molecular association in the case of electrolytes appear therefore as parts of a general phenomenon; they are complementary, the precise molecular condition being a matter almost wholly of the dielectric character of the solvent.

The point at which, in the dielectric constant list, molecular association begins to be apparent cannot well be decided because the molecular association depends on the concentration also; but in solutions which are not really concentrated, association is clearly in evidence for a number of salts, such as tetrapropylammonium iodide and quinoline ethiodide ¹⁵² in alcohol, and in acetone the indication is still clearer. As the dielectric constant of alcohol is 180 at its boiling point, this value may be considered as the one below which molecular association begins to be evident. Nor is this the case with salts alone; for amides and anilides show a tendency, except in quite dilute solution, to association in alcohol and acetone. With salts, except chlorides, in a few cases, a dielectric constant above 180 leads to a molecular weight below the normal.

Finally, it may be pointed out that the associated condition of a solvent does not endow it with dissociating power, unless, at the same time, its dielectric constant is high. Thus, water, formic acid, formamide, acetonitrile (except with chlorides) and ethylene cyanide are associated substances which do produce dissociation of salt molecules; but dimethylethylcarbinol, phenol, acetic acid, and *iso*-amyl alcohol are also associated yet permit pronounced association. The first five have high dielectric constants, however, the remainder only low values. For weak electrolytes and non-electrolytes this is likewise true. Discounting association in water, both benzoic acid and acetic are strongly associated in liquid nitrogen peroxide and the former also in ethylene cyanide (see p. 43).

CHAPTER V.

MOLECULAR COMPLEXITY IN THE LIQUID STATE.

THE determination of the molecular condition of matter in any condition other than the diluted state is attended at present with very considerable difficulties. Avogadro's hypothesis makes determinations in the case of gases and dilute solutions a very simple matter in theory, but for liquids there is at the present time no similar guiding principle. With one or two exceptions, indeed, all the methods available are empirical.

The basis of most methods, therefore, consists in instituting a comparison of the molecular properties of different liquids. As the outcome of such tests, it is found that the molecular property depends very largely on the molecular weight of the substance; that is to say, the property is additive. As such, it is possible to express it in simple mathematical form as an equation involving the molecular weight and a constant. Thus, Trouton's Rule (see p. 52) is the expression of the experimental fact, that the heat required to evaporate a molecular weight in grams of a given liquid is proportional to the absolute temperature of the boiling-point, or

Ml = kT.

Such generalizations on molecular properties are almost always subject to exceptions, so that it is possible to divide liquids into two classes, those which obey the rule, and those for which k is greater or less than the usual value. The value of M adopted in all expressions such as the above is the molecular weight which the substance has in the state of vapour. Whether M is truly the molecular weight of the liquid will be discussed later, but this much can be said, namely, that for the large body of liquids which obey the given equation, the degree of complexity must be the same, and either, on condensation from the gaseous state, no change of molecular size occurs, or, a change

4

in complexity in every case the same occurs yielding an aggregated molecule which undergoes no variation in size in the liquid state. Such substances are commonly spoken of as normal.

The most obvious explanation to account for the second or abnormal class, is to suppose that the molecular weight is not truly represented by M, but that it is either greater or smaller, and is represented by xM, where x represents the degree of complexity, which can be determined knowing the other quantities in the equation. This explanation is rendered very probable because it is found that substances associated in solution, are always abnormal in the liquid condition. Hence, it is but a natural conclusion to assume that the real molecular weight is not M but xM, where x > 1. Both qualitative and quantitative methods of determining association follow such lines. In the present chapter certain qualitative evidence of association will be given, but it is easy to discern that many of the methods quoted could, without difficulty, be extended also to a quantitative determination.

Evidence of Molecular Association in Liquids.

I. From Vapour Pressure Data.—It is a very common occurrence that a substance which is volatile possesses in solution the normal or simple molecular weight, and it is only a step to the deduction that association of a liquid will diminish its vapour pressure. Thus, one finds that sulphuric acid at ordinary temperatures possesses a scarcely measurable vapour tension, a fact which receives ready explanation from its associated state.¹⁵⁴

The effect of molecular association on the vapour pressure comes out clearest when tested by the expression of van der Waals, 155 log p_c – log $p = k \frac{T_c - T}{T}$, by which the vapour pressure of a liquid can be calculated from the critical temperature and pressure, T_c and p_c , and the temperature T for which p is required. The constant k was found by Young 156 to have a mean value of 3.06 for a large number of substances; but for hydroxylic substances the factor k is not constant and always exceeds 3.06, as the following results indicate:—

Water . . . 3 '20-3 '24 Ethyl alcohol . . 3 '58-4 '02 Methyl alcohol . . 3 '56-3 '77 Propyl alcohol . . 3 '49-3 '77 Acetic acid 3 '36-3 '49

The fact that these substances in solution can exist as complex molecules renders an explanation of these abnormal results as due to association an obvious one.

2. Boiling-point Evidence.—(a) Just as the vapour pressure may be diminished by association, so the boiling-point will be raised. Now the boiling-point, although not strictly an additive property, has been demonstrated to be very largely influenced by the mass of the molecule, so that it is possible to calculate the boiling-point, very approximately, of a given substance, knowing its chemical composition, by adding the values for the elements. In this way the value calculated by van't Hoff for water is - 207° instead of + 100°. 157

Walden 158 proceeds in a somewhat different manner. Methyl sulphide (B.-pt. 37°) has a boiling-point 60° higher than methyl ether (B.-pt. - 23°), whilst the difference between ethyl sulphide and ethyl ether is 58.4°. Substitution of oxygen by sulphur, therefore, raises the boiling-point, on the average, 59°. Compared with sulphuretted hydrogen (B.-pt. - 61°) water therefore should boil at - 120°. The actual boiling-point, 100°, can readily be accounted for if liquid water is associated.

(b) Another way in which the effect on the boiling-point can be indicated is by means of the equation

$$\frac{\mathbf{T'_A}}{\mathbf{T'_B}} = \frac{\mathbf{T_A}}{\mathbf{T_B}} + \mathbf{C} (t' - t)$$

relating the boiling-points under different pressures of two substances A and B, TA and TB being the respective boiling-points under the same pressure, T'A and T'B the boiling-points under some other common pressure, and t' and t the boiling-points of either of the substances under the given pressures, and c a constant. When A and B are very closely alike in properties, as, for example, chlorobenzene and bromobenzene, c becomes exceedingly small. In any case, the equation for most substances corresponds to a straight line, but for alcohols and aliphatic acids it does not. 159 Again, the deviation may be explained by assuming the associated condition of these hydroxylic substances.

3. Latent Heat.—Closely connected with the boiling-point is the latent heat absorbed during the passage from liquid to vapour. Unless there is a difference between the molecular state of the liquid and the vapour, the heat imparted will be employed in

performing external work. A dissociation into a simpler molecular state will call for an increased supply of heat, that is to say, the latent heat of associated substances tends to be high. An explanation, depending on association, has also been suggested for the maxima observed in the latent heat of some liquids, such as with ethyl alcohol, the latent heat of which reaches a maximum at 10° and acetic acid at 120°.160

Trouton's Rule enables one to test the molecular state of a substance by a measurement of its latent heat. As previously stated, the mathematical expression of the rule is

$$\frac{Ml}{T}$$
 = constant.

Although strictly the equation should apply only to similarly constituted liquids, as it is only for similar substances that corresponding states can be realized,* it is found in practice to hold for a large number of substances. An investigation by Walden 161 has proved the average value of the constant to be 20.7, and that departure from this rule, either by a higher or a lower value, may be taken as evidence of molecular association. Thus, in the following table, are both types of deviation from k = 20.7.

Substance.	$\frac{\mathbf{M}l}{\mathbf{T}}$.	Substance.	Ml T
Water	25.9	Acetonitrile	17.74
Formic acid	14.8	Propionitrile	19.97
Acetic acid	14.88	Butyronitrile	20.33
Methyl alcohol	24.67	Valeronitrile	19.89
Ethyl alcohol	28.3	Acetone	22.2
Phenol	28.2		

The high numbers appear to be obtained with hydroxylic substances, low values with organic acids and nitriles.

It might be thought that the method should be capable of use in determining the molecular weight, or at any rate that the extent of deviation from the mean value for normal substances should indicate the extent of association, but comparison with methods given in the next two chapters will show that it does not serve such a purpose. Phenol, for example, deviates much more than either water or methyl alcohol although less associated than either. In this connexion what has already been said about the comparison being limited to substances of similar

^{*} For the derivation of the equation thermodynamically see van't Hoff, "Lectures on Theoretical and Physical Chemistry," Part III, p. 57.

53

constitution must be borne in mind. Nernst 162 has found that the value of $\frac{Ml}{T}$ depends on the boiling-point of the substance, and increases as the boiling-point rises, so that for hydrogen the ratio is only 12.2. Further, he has shown that

$$\frac{Ml}{T} = 9.5 \log T - 0.007 T$$

when the substance is normal, but when associated,

$$\frac{Ml}{T}$$
>9.5 log T - 0.007 T.

Thus, for water and ethyl alcohol the ordinary expression gives values 25.9 and 26.9 (quoted by Nernst) respectively, whilst the new equivalent leads more to the normal value, 21.8 for water, and 21.7 for alcohol being the actual figures. The low value for acetic acid, he suggests, occurs because the vapour molecules are also associated, which is not the case with alcohol. When the actual molecular weight, 97, in the state of vapour is employed for M, then $\frac{Ml}{T}$ becomes 22.1. For acetic acid the value of 9.5 log T – 0.007 T which, it will have been noticed, does not contain the molecular weight as a term, amounts to 21.9.

Although Nernst's explanation may apply to acetic acid, it does not appear to apply to the nitriles, for experiments on acetonitrile vapour in the writer's laboratory do not indicate any appreciable association, like acetic acid, and Trouton's Rule must therefore be considered at best as only affording qualitative evidence of association (cp. also application of method in chap. VII.).

Traube ¹⁶³ has applied the Rule to test the molecular state of bismuth, iodine, sulphur, zinc, cadmium, mercury, and bromine, calculating the latent heat from the thermodynamic formula

$$l = T \cdot \frac{dp}{dt} (v_1 - v_2).$$

The normal value of Trouton's Constant was found in all cases except sulphur, where it had the value 30.2. For the metals the monatomic state appeared to be the normal liquid condition.

- 4. Evidence from Critical Data.—
- (a) Divergence from the Rectilinear Diameter.—The now well-known rule of Cailletet and Mathias for the determination of the critical volume of a substance is based on the observation that

the mean values of the densities of the saturated vapour and the liquid with which it is in contact lie on a straight line (the rectilinear diameter) passing through the critical point. Subsequent investigations ¹⁵⁹ have proved that the rectilinear diameter has in a number of cases a very slight curvature which can be readily corrected, but not for alcohols and water. Here the curvature is quite distinct. The inference is that the liquid and vapour in contact have different degrees of complexity. Acetic acid, however, gives a diameter nearly straight, as with this substance both the liquid and the vapour are associated.

(b) Irregularities in the Critical Volume.—The determination of critical volume provides a second test of molecular association; for Young and Thomas ¹⁶⁴ discovered that most liquids have a density at the critical temperature of 3.85 times the value calculated on the basis of the gas laws. Once again the alcohols and acetic acid are abnormal, their factors being

Methyl alcohol	4.2
Ethyl alcohol	4.03
Propyl alcohol	4.02
Acetic acid	5.00

(c) Evidence from the Value of the Critical Coefficient.— Utilizing as a basis the characteristic equation of van der Waals applied at the critical point, namely,

$$\frac{3}{8}\left(\mathbf{I} + \alpha\theta\right) = \frac{\pi\phi}{\left(\mathbf{I} + 3\pi\phi^2\right)\left(\mathbf{I} - \frac{\phi}{3}\right)},$$

together with Maxwell's electro-magnetic theory of light and Clausius' equation connecting the dielectric constant with the ratio of the actual to the apparent volume of the molecules of a substance, Guye ¹⁶⁵ obtained a relationship

$$\frac{273 + \theta}{\pi} = \frac{I}{f}. MR,$$

where $(273 + \theta)$ represents the absolute critical temperature (T_c) π the critical pressure (p_c) , MR the molecular refraction and f a constant. Now the value of f which a large number of normal substances have is 1.8, but for water, methyl alcohol, and acetic acid it amounts only to about 1.1.

It will be clear that since the equation contains the value of the molecular weight M, it should be possible to determine the molecular complexity of an associated substance. This has been done. Unfortunately, the method is limited to the critical temperature and therefore also involves practical operations which are not easy to perform.

Other evidence of association is forthcoming from a study of corresponding states. At a given reduced pressure $\frac{p}{p_c}$, where p is the pressure employed and p_c the critical value, the ratios $\frac{T}{T_c}$, $\frac{v_l}{v_c}$ and $\frac{v_v}{v_c}$, where T_c and v_c are the critical temperature and volume, respectively, T the temperature at pressure p, and v_l and v_l the corresponding volumes of liquid and vapour, should be constant for all substances. Although not perfectly true for normal substances, the divergencies in the case of the alcohols and acetic acid are much greater.

5. Evidence from Viscosity Measurements.—If the viscosity of a series of substances is tested, it is found to increase in value as the molecular weight rises. A high viscosity, therefore, indicates usually a high molecular weight; so that if the simplest molecular weight of the substance examined is low, the inference to be drawn from a high viscosity is that association of molecules has occurred. In the main this is the argument on which the viscosity test of association rests. Thus, Thorpe and Rodger, from the rather high and irregular viscosities of the alcohols and aliphatic acids, supposed association to occur.

There are several methods, mainly the work of Dunstan and his collaborators, in particular Thole, of demonstrating the existence of association by means of viscosity data. One method * depends on the discovery that the expression $\frac{\eta}{Mv} \times 10^6$, where η is the viscosity coefficient and Mv the molecular volume is a constant quantity for series of like substances. Thus:—

Substance.	$\frac{\eta}{\mathrm{M}v} \times 10^6$.	Substance.	$\frac{\eta}{Mv} \times 10^6$.
Benzene	65	Alkyl chlorides	37.4
Chloroform	67	Alkyl bromides	50.7
Carbon disulphide	бо	Alkyl iodides	68.3
Ethers	25'2	Ketones	43'3

All the substances contained in this table are normal. When

^{*} For a fuller account of the effect of association on viscosity, reference should be made to Dunstan and Thole's "The Viscosity of Liquids" in this series,

association occurs, the value of the expression is much higher, as in the following examples:—

Substance.	$\frac{\eta}{Mv} \times 10^6$.	Substance.	$\frac{\eta}{\mathrm{M}v} \times 10^6$.
	Mv		
Water	493	Formamide	181
Methyl alcohol	138	Acetamide	219
Ethyl alcohol	189	Propionamide	162
Acetic acid	195	Urethane	103
Glycerine	106,000		

A variety of other methods, especially in the case of water, have been proposed for the detection of association in liquids, and will be referred to in chapters VII. and VIII.

CHAPTER VI.

SURFACE TENSION METHODS OF DETERMINING MOLECULAR COMPLEXITY IN THE LIQUID STATE.

SURFACE tension methods of investigating the molecular condition of liquids, methods which have received far greater attention than any other, may be divided roughly into two classes according as they are related to the Ramsay and Shield's method and depend on the measurement of surface energy, or as they are based on a study of the specific cohesion, as in the methods of Kistiakowsky, Walden, Dutoit and Moijou.

The method of Ramsay and Shields ¹⁶⁶ which has exerted so great an influence on other methods, and was indeed the first by which insight was obtained into the molecular state of liquids, depends on a study of the temperature coefficient of molecular surface energy, as expressed in the equation

$$\gamma (M v)^{\frac{2}{3}} = k (\Gamma_c - t - d^\circ) = k (\tau - d)$$

in which γ , M, and v represent the surface tension, molecular weight, and specific volume of the substance at the temperature τ measured from the critical temperature (T_c) downwards, and k is a constant.

The above equation differed from that of Eotvos, ¹⁶⁷ whose work underlay that of Ramsay and Shields, in the factor d° , usually 5° or 6°, a correction necessary because the surface tension varies proportionately with the temperature only from some point a little way below the critical. In defining the molecular surface as $(Mv)_{3}^{2}$, Ramsay and Shields followed Eötvos, assuming that the molecules at the liquid surface are distributed in the same manner as throughout the rest of the liquid. For this assumption, they stated explicitly that there was no direct evidence, but that the results of the method appeared to make the assumption justifiable. It is possible that the abnormal results found later may be due to this assumption not being wholly true. The

value of k,

$$k = \frac{\gamma (Mv)^{\frac{2}{3}}}{\tau - d} = \frac{\Delta E_o}{\Delta t}$$
 (a shorter expression sometimes employed)

was found for a number of liquids which could reasonably be assumed as normal, to have the mean value of 2·121 over a range of temperature from about – 90° up to the critical point.

The equation $\gamma (Mv)^{\frac{3}{3}} = k (\tau - d)$ has an analogy to the gas equation PV = RT, in that in the former the molecular surface energy is expressed as proportional to the temperature measured from the critical point, in the latter the molecular volume energy as proportional to the absolute temperature; and just as association or dissociation of the molecules of a gas affect V or P and will be reflected in the value of R, so, in like manner for a liquid, a variation in the value of k should indicate similar changes.

Suppose, then, that a liquid is abnormal. If the requisite measurements have been carried out, the value of the factor in the equation will no longer be k but some other value, k_1 , if the numerical value of the molecular weight employed is the normal value, M. The equation thus becomes

$$\gamma(Mv)^{\frac{2}{3}} = k_1(\tau - d)$$
 . (1)

By inserting the true value of the molecular weight, the factor will assume its normal value, namely, $k=2\cdot12$. Let this true molecular weight be xM, where x is the degree of complexity. Then

$$\gamma(xMv)^2 = k(\tau - d) \quad . \qquad . \qquad . \qquad (2)$$

By comparing (1) and (2)

$$x = \left(\frac{k}{k_1}\right)^{\frac{3}{2}} = \left(\frac{2 \cdot 12}{k_1}\right)^{\frac{3}{2}}$$

The determination of the factor d would make the method rather difficult, but the necessity of evaluating d can be removed by making measurements at two different temperatures, when the following relations hold:—

$$\gamma_1 \left(\mathbf{M} v_1 \right)^{\frac{2}{3}} = k \left(\tau_1 - d \right)$$

 $\gamma_2(Mv_2)^{\frac{2}{3}} = k(\tau_2 - d)$, neglecting any molecular change between τ_1 and τ_2 , so that M and k remain constant.

By subtraction

$$\frac{\gamma_1 (M v_1)^{\frac{2}{3}} - \gamma_2 (M v_2)^{\frac{9}{3}}}{\tau_1 - \tau_2} = k$$

and $\tau_1 - \tau_2$ is merely a temperature interval independent of any scale. A second method is to write the equation in the differen-

tial form (see p. 61). The practical measurements involved are those of surface tension and density.¹⁶⁸*

The following table drawn from different sources is representative of different types of substances and will indicate the degree to which liquid substances appear to be associated.

Substance.	Temperature.	k.	x.
(Water	0°-140°	0.84 -1.51	3.81-2.32
Methyl alcohol	16·0°- 78·0°	0.933-0.969	3'43-3'24
Ethyl alcohol	16.0°- 78.0°	1.083-1.12	2.74-2.43
Acetic acid	16.0°-132.0°	0.000-1.024	3.62-2.77
{ Acetone	16.8°- 78.3°	1.818	1.36
Phosphorus (P ₄)	78·3°-132·1°	2.202	0.94
Nitrogen peroxide (N2O4)	1.6°- 19.8°	2.110	1.01
Phenol	46.0°-184.0°	1.682-1.899	1.42-1.18
Nitroethane 169	16.6°- 79.6°	1.647-1.684	1.46-1.41
o-Chlorophenol	12.7°- 73.3°	2.50 -5.00	1.0
p-Chlorophenol	51.6°- 99.8°	1.86 -2.03	1.55-1.08
{o-Cresol	39.6°- 99.7°	1'97 -2'04	1.13-1.09
m-Cresol	19.1 _o - 99.9 _o	1.63 -1.73	1.48-1.33
p-Cresol 170	45'5°- 99'8°	1.24 -1.85	1.62-1.56
n-Propylamine	10°- 45°	1.84	1.24
iso-Amylamine	20°- 60°	1.955-2.012	1.13-1.08
Dimethylnitrosoamine	20°- 75°	1.42-1.942	1.34-1.14
Formamide	20°- 75°	0.63 -0.95	6.18-3.34
Acetamide	85°-120°	1.19 -1.50	2.47-5.11
Lactamide	80°-120°	1.02 -1.14	2.87-2.54
Benzamide	130°-170°	1.002-1.43	2.70-1.81
Formanilide	60°-105°	1.485-1.725	1.69-1.36
Ethyl urethane 171	60°-105°	1.492-1.22	1.69-1.22
(Acetonitrile 172	20°- 60°	1.42 -1.26	1.43-1.28
Lactonitrile 171	20°- 60°	1.255-1.325	2.51-5.03
Propionitrile 173	20°- 60°	1.63 -1.66	1.48-1.44
(Sodium nitrate	330°-400°	0.445	10.68
Potassium nitrate 174	341°-407°	0.203	8.73
(Silver chloride	446°-580°	0.959	3.50
Lead chloride 175	480°-600°	0.003	3.60
(4	- 303	5 00

Criticism of the Ramsay and Shields' Method.

Subsequent work has considerably reduced the value of the Ramsay and Shields' method. The criticisms which have been levelled at it as a means of actually measuring the degree of association are (I) that the equation requires modification, (2) that the value of the factor k, 2·12, is not always valid, (3) a much more sweeping criticism, that surface tension methods of measuring the molecular complexity do not afford a true indication of the molecular state of the liquid as a whole.

^{*} For apparatus and details of measurement see Ramsay and Shields, Hewitt and Winmill, Turner and Merry, Walden, Dutoit and Moijou. 168

The need of a more accurate method of determining the value of x was recognized by Ramsay himself, and before the appearance of other criticism, he revised the original expression and employed another deduced by Rose-Innes, 160 namely,

$$\gamma (Mv)^{\frac{2}{3}} = \frac{k_1 (\tau - d)}{1 + \mu \tau},$$

where μ and k_1 are new constants to be derived for the substance. This formula was found to represent the variation of the molecular surface energy of water, the alcohols and acetic acid, over a range of temperature from 30° below the critical point down to low temperatures.*

Now, as previously stated, the original equation for associated liquids becomes

$$\gamma (xMv)^{\frac{2}{3}} = k (\tau - d)$$
, where $k = 2.121$.

Dividing this by the Rose-Innes equation,

$$x = \left[\frac{2 \cdot 1 \cdot 2 \cdot 1}{k_1} (1 + \mu \tau)\right]^{3/2}$$

When applied by Ramsay to the earlier experimental data, the following numbers were obtained:—

Substance.	Temperature.	k_1 .	μ.	x.
Water	0°-140°	2.631	0.00318	1.707-1.589
Methyl alcohol	-89.8°-220°	1.489	0.00104	2.65 -1.75
Ethyl alcohol	-89.8°-230°	2.170	0.00103	2.03 -1.00
Acetic acid	20°0°-280°	1.010	0.00163	2.13 -1.30

A noticeable reduction in the value of the degree of association is brought about by the new formula. Indeed, Tyrer argues that the results now obtained are too low, the original equation probably giving more correct results, because the value of the critical temperature required in the new formula depends on the degree of complexity and will accordingly be variable (see more fully, Batschinski, p. 65). The real source of error in the original equation, apart from the principle, lies, according to Tyrer, in neglecting the variation of molecular complexity with temperature over the range for which k is found. For the equation

$$\gamma \left(\mathbf{M} v \right)^{\frac{2}{3}} = \frac{k_1 \tau - k_1 d \left(\mathbf{I} - \mathbf{IO}^{-\lambda \tau} \right)}{\mathbf{I} + \mu \tau},$$

where λ is a new constant. A somewhat similar formula is that of van der Waals, 176 namely,

$$x^{\frac{2}{3}} = \frac{k \left(\tau_1 - \tau\right) + \gamma \left(Mv\right)^{\frac{2}{3}}}{\gamma \left(Mv\right)^{\frac{2}{3}}}.$$

^{*} A still more accurate expression 160 is

 $\gamma(xMv)^{\frac{2}{3}} = k \left(T_c - t - d \right), \text{ written in the differential form becomes}$ $x^{\frac{2}{3}} \frac{d}{dt} \gamma \left(Mv \right)^{\frac{2}{3}} + \gamma \left(Mv \right)^{\frac{2}{3}} \frac{d}{dt} x^{\frac{2}{3}} = k \frac{dT_c}{dt} + k.$

The second term was taken into account in the revised formulæ used by Ramsay, but the third was neglected as being too small. In practice it cannot always be neglected.

The second cause for criticism arises from the fact that the value of k, originally thought to lie very closely in the neighbourhood of 2·12, may, even for non-associated substances, vary widely from this value. Even Ramsay and Aston ¹⁷⁸ noticed that several esters had values of k between 2·2 and 2·3, but did not study the ester series sufficiently to discover the abnormalities which subsequent investigators brought to light. Thus, the higher esters of malic acid, ¹⁷⁹ amines, which show in a very clear way the effect of substitution and increase in the molecular weight on the value of k, phenylurethane and the alkylated anilides, ¹⁷¹ tristearine and tripalmitine, ¹⁸⁰ and many other substances, ¹⁸¹ have values of k much higher than 2·12, as the following table shows:—

Temperature.	k.
12.3°-102.0°	3.68 -3.38
16.3°-106.6°	3.20 -3.46
54.6°-150.0°	3*34 -3*33
20°- 75°	2.08 -2.225
20°- 75°	2.805-3.00
95°-135°	3.41 -3.58
20°- 60°	1.955-2.015
20°- 75°	3.102-3.182
57°7°-134°7°	6.51 -2.32
55'7°-125'1°	4.92 -4.96
	12'3°-105'0° 16'3°-106'6° 54'6°-150'0° 20°- 75° 20°- 75° 20°- 60° 20°- 75° 57'7°-134'7°

If the mode of interpretation set out by Ramsay and Shields is to be strictly followed, then the above substances must be regarded as undergoing dissociation in the liquid state, to such an extent that tribenzylamine is split into two parts (x = 0.49 - 0.47) and tripalmitine into four or five (x = 0.203 - 0.240). But there are many cogent reasons why this is improbable; for molecular weight determinations in several solvents give no indication of the extensive association suggested from the determination of $k^{171, 180}$. Moreover, the coefficients of surface energy of tristearine and tripalmitine in benzene solution have nearly the normal value, Trouton's Constant for the esters tabulated has

the normal value, ¹⁸² and several other methods of determining the molecular complexity of liquids one and all afford evidence against the hypothesis of dissociation. Homfray and Guye, ¹⁷⁹ however, favoured the hypothesis of dissociation and, more recently, Guye¹⁸³ has put forward the view that the esters are dissociated into hydrocarbon and acid by reason of the powerful forces acting on the surface film of a liquid, in which part alone, he considers, the dissociation takes place. On this ground, he regards the Ramsay and Shields method as unsatisfactory, since the molecular state of the liquid at the surface and in the interior will be different.

The whole subject of the variation of the temperature coefficient, k, has been systematically attacked by Walden and Swinne, who find that its value is really a function of molecular constitution. Thus, whilst certain groups tend to produce association, the values of $\frac{\Delta E}{\Delta t} = k$, as also the temperature coefficient of

molecular cohesion, $\frac{\Delta \ \mathrm{M} \, a^2}{\Delta t}$ (vide, p. 71) are affected by substitu-

tion. An increase in the value of the coefficient accompanies, for example, substitution of CH_3 by C_6H_5 , of O by S, N by P or Sb, Cl by Br, and Br by I in a measure approximately quantitative, whilst a steady rise always occurs in a homologous series. As in every case, these changes are accompanied by an increase in mass, it is obvious that, despite minor constitutive influences, the temperature coefficients, both of molecular surface energy and molecular cohesion are additive properties. Two striking facts give proof of this. The first is that tripalmitine and tristearine, the substances of highest molecular weight so far investigated, have also the greatest temperature coefficients; the second, and still more conclusive, that the additive character can be expressed by the empirical formulæ

$$\frac{\Delta E}{\Delta t} = -\left[1.90 + 0.011 \left(\Sigma /A\right)\right]$$

$$\frac{\Delta M a^2}{\Delta t} = -\left[0.00027 \left(\Sigma /A\right) + 0.0103\right] \text{ where } t = 100^\circ,$$

and $\Sigma \sqrt{A}$ represents the sum of the square roots of the atomic weights of the constituent elements, and by means of these formulæ the values of the coefficients can be calculated for many substances of widely different character and molecular weight.

The great differences possible in the molecular weights of

different liquids may be attended by another consequence, namely, a wide variation of critical temperature. Now in order to compare accurately the values of the molecular surface energy, as of any other physical property, the comparison ought strictly to be carried out at corresponding temperatures, a condition which in practice it is most difficult to attain. The manner in which k varies, decreasing with rise of temperature as with tristearine and tripalmitine suggested the possibility that at the critical temperature, at which all substances are in corresponding conditions, normal values might be obtained. In order to test this possibility, Walden and Swinne calculated the values of the critical temperatures by the aid of the Batschinski formula (p. 66) and on extrapolating the value of k to this temperature, found it to have the value 2'II for tripalmitine and 2'06 for tristearine. While these two substances lend most favourable support to the suggestion, a number of others are known for which the coefficients rise with the temperature, so that the normal value at the critical point could not, with good reason, be predicted.

As to the dissociation hypothesis of Guye, Walden and Swinne brought forward several arguments which were not in harmony with it. But the most obvious arguments, in the writer's opinion, lie in the additive character of the coefficients, and in the widely differing classes of substances showing these high values, substances the mode of dissociation of which it would be difficult to surmise.

To sum up the evidence of this section, Ramsay and Shields' method leads in some cases to low values of k, values which can with confidence be explained as due to molecular association; in some other cases, now known to exceed in number the original normal examples of Ramsay and Shields, high values are obtained which cannot be explained by the reverse process of dissociation. Against such a supposition, the results of other methods of determining the molecular complexity of liquids, for example, Trouton's Rule, already dealt with, Walden's, Kistiakowsky's and that of Dutoit and Moijou all yet to be discussed, are in agreement. As the deduction of the degree of association by Ramsay and Shields' method involves the use of a factor k which varies widely even for normal substances, it is obvious that the true molecular complexity cannot be calculated. There is the possibility, however, especially in view of the additive char-

acter of k of limiting the method to substances having molecular weights within a certain range, dividing substances, that is to say, into groups having molecular weights within a certain range and finding the value of k for such a restricted group, that quantitative results expressing at least the relative order of complexity might be obtained.

Three other methods, related to Ramsay and Shields' method, will now be described. Two of them seek to correct the original method, the third, however, deviates considerably.

Morgan's Method.

This method ¹⁸⁴ is really only qualitative, enabling one merely to discover the existence of association; but it is more appropriately described here than in the preceding chapter. The author of the method believes that some of the difficulties arising from the use of the Ramsay and Shields' method are due to the mode of measuring the surface tension.

In the course of experiments on the weight of a falling drop formed at and falling from the end of a tube under no force save that of gravity, Tate ¹⁸⁵ found that the weight of the drop was proportional to the weight which would be raised in the same tube by the force of capillarity; that is to say, the weight of the falling drop is proportional to the surface tension.*

It is this fact, tested by Morgan and his co-workers for liquids other than water, which forms the basis of the method.

Supposing w to be the weight of a drop formed under the conditions stated, then w is proportional to γ , the surface tension, and the Ramsay and Shields' equation may be written

$$w\left(\frac{M}{d}\right)^{\frac{2}{3}} = K_B (t_c - t - a)$$

where d is the density of the liquid, a corresponds to d° in the original equation, t_c and t are the critical temperature of the substance and the temperature of experiment, respectively, and K_B is a constant for the particular glass tip at which the drop is formed. To standardize the apparatus and determine K_B , benzene, of critical temperature 288·5°, is employed, and the value of w determined for drops falling from capillary tubes bevelled at the end at an angle of 45°, so as to give tips of from 4·5 mm. to 5·5 mm. diameter.

^{*} Lohnstein 196 however, argues that this relation is, in general, not true.

If desired, the surface tension of the liquid can be deduced from the formula $\frac{wk}{K_B}$, where k is the Ramsay and Shields' constant and K_B that in Morgan's formula. In this way, it is maintained, more consistent measurements of the surface tension can be made than by the method of capillarity or any other method.

It is unnecessary, however, to find the value of γ in order to determine whether a liquid is or is not associated. The test applied is to calculate the value of t_c in the above equation. Should the value be independent of the temperature of measurement, the liquid is regarded as non-associated, and in this case, from the experimentally determined value of t_c , M, the molecular weight, can in turn be measured.

Supposing, however, the value of t_c varies with the temperature throughout the series of measurements, then the substance is regarded as associated. By this test, the alcohols and aliphatic acids, formamide, and several ketones have been shown to have higher molecular weights in the liquid than in the gaseous state. No attempt has been made, however, to determine the actual degree of complexity, or to attack the real difficulties arising from the surface tension method.

Batschinski's Method.

While the preceding method lays stress on the actual operation by which surface tension is to be measured, Batschinski 187 accepts existing experimental data but modifies the original equation. In doing so he introduces an interesting idea that the associated and non-associated forms of any one substance must be treated, from the physical point of view, as distinct individual substances. The three constants R, a and b of van der Waals' equation, which are characteristic of the individual substance, must be different for the single and complex molecules. Whilst this is true of R and a, b is held to be less dependent on association and may, indeed, be considered as approximately independent 187a . Therefore, of the critical constants, temperature, pressure and volume, viz.

$$T_c = \frac{8a}{27 \text{ Rb'}}, p_c = \frac{a}{27b^2} \text{ and } v_c = \frac{I}{3b'}$$

only v_c will be the same for the associated and simple molecules, whereas T_c and p_c will be functions of the condition of the substance. A complex molecule should therefore have a differ-

ent critical temperature and pressure from the single molecular form, provided its degree of complexity could be retained unchanged over the temperature range up to the critical point; but as, in general, the degree of complexity varies with the temperature, so also will the critical, or meta-critical values (T_{mc}, P_{mc}) as Batschinski calls them, alter. If the dissociation of the molecule is complete, only then does the meta-critical become identical with the critical temperature.

Now in the Ramsay and Shields' formula for associated substances,

$$\gamma (x Mv)^{2/3} = k (T_c - t - d) = k (\tau - d),$$

the value of T_c must be substituted by T_{mc} .

To deduce the value of T_{mc} , the variation of some property must be studied at those temperatures for which the surface tension is also measured, or calculated from existing data at other temperatures by known formulæ. The property made use of is the viscosity, in connexion with which Batschinski ¹⁸⁸ had previously deduced the relationships

$$\eta T^3 = \text{constant, and}$$

$$\frac{T d_c^{\frac{1}{r}}}{(\eta T^3)_r^2} = \text{constant,}$$

where d_c and T_c are the critical density and temperature, respectively, η the viscosity and T the absolute temperature of its measurement.

The mean of ten closely agreeing values of this constant, using non-associated substances, hydrocarbons, esters and ether, was found to be 16.31. For associated substances, however, T_c must be substituted by T_{mc} , when it is assumed that

$$\frac{T_{mc}d_c^{\frac{1}{7}}}{(\eta T^3)^{\frac{2}{7}}} = 16.31$$

and in turn, that the meta-critical temperature can be calculated by using the value of the constant given. If d_c is unknown, then an alternative, but somewhat less accurate, expression is used,

$$\frac{T_{mc}d_0^{\frac{1}{7}}}{(\eta T^3)_T^2} = \text{constant} = 19.4,$$

where d_0 is the density at 0°.

Substituting T_c by T_{mc} in the equation of Ramsay and Shields, the values of x may be found, and Batschinski obtained results from which the following is a selection:—

Substance.	Temperature.	Tmc.	<i>x</i> .
Water	0°-100°	752.3°-581.4°	2.83-1.10
Methyl alcohol	20°	590°1°	3.43
Ethyl alcohol	20°-60°	719 [.] 9°-657 [.] 6°	4.43-3.65
n-Propyl alcohol	16·4°-78·3°	878·1°-707·4°	5.03-3.12
n-Butyl alcohol	17'4°-77'9°	959 · 4°-706 · 7°	4.79-3.03
Formic acid	16·8°-79·4°	783·1°-690·6°	3.80-3.68
Acetic acid	20°0°-100°0°	700°0°-650°6°	3'47-2'74
Propionic acid	16·6°-132·5°	698·9°-656·6°	2.48-5.26

The results of this method are on the whole to enhance the values of the degree of complexity. At the same time it will be noticed that the most associated member of the alcohol series is not methyl alcohol, as usually found, but that association increases with molecular weight up to propyl alcohol.

Bennett and Mitchell's Method.

The total surface energy, λ , of a liquid, given by thermodynamics as

$$\lambda = \gamma - T \frac{d\gamma}{dT},$$

where γ is the surface tension, has been shown by Whitaker ¹⁸⁹ to be of more importance for the study of the physical properties of a liquid than the surface tension.

This same quantity, λ , has been connected with the equation of Ramsay and Shields by the following expression due to Kleemann ¹⁹⁰

$$\left(\gamma - t \frac{d\gamma}{dt}\right) (Mv)^{2/3} = 2.12 (T_c - 6).$$

Bennett and Mitchell's method ¹⁹¹ is based on this result. In the first place, the temperature variation of the quantity

$$\left(\gamma - t \frac{d\gamma}{dt}\right) (Mv)^{2/3}$$
 which is the total molecular surface energy, is

only expressed approximately when the numerical constant is 2·12, and Bennett and Mitchell found a more precise relationship

$$\left(\gamma - t \frac{d\gamma}{dt}\right) (Mv)^{\frac{2}{3}} = 2.38 (T_c - 6).$$

Here, then, is an expression very similar in form to the equation of Ramsay and Shields, the difference being that the total surface energy is the fundamental property instead of the surface tension, and the introduction of this quantity slightly alters the constant of the equation.

The total molecular surface energy is written as K, and was

proved to remain constant over a wide range of temperature for each non-associated liquid examined. Moreover, it was found to be additive in character in such a way that values could be assigned to the atoms as indicating their contribution to the total molecular surface energy; so that when K had been determined for a number of normal compounds, atomic surface energies could be determined like atomic volumes or refractions are deduced.

Atomic (or Group) Surface Energies.

CH ₂	83	C_6H_5	990	NOH	820
H	310 .	CN	870	NO_2	700
С	- 537	CNS	945	Double bond	654
O (in ether)	160	NCS	465	Ring closing:-	
O (ketonic)	78o	C1	600	(1) Hexamethylene	740
N (in NH ₂)	0	\mathbf{Br}	805	(2) Pentamethylene	717
		I	820	(3) Trimethylene	671

From these values, K can be calculated with fair accuracy for a large number of compounds, only the esters of acylmalic acid and of polybasic acids deviating to any considerable extent, and, save for the esters mentioned, the substances which give an abnormally high value of Ramsay and Shields' constant are normal when subjected to this new test.

From these regularities associated substances are excluded and marked deviation may be employed as a test of association. The actual degree of complexity of associated substances is determined by a study of K, which, although found to be less than the normal value calculated from the atoms, rises regularly with the temperature. Assuming this rise to be due to a continuous breaking down of complex molecules, a constant value may be expected to be arrived at eventually when the substance is no longer associated. This limiting value of K, written as K', should, when extrapolated from the data at lower temperatures, equal the number calculated for the substance from the atomic values. The importance of being able to calculate K is therefore obvious. For a non-associated substance, there are thus two equations,

$$\left(\gamma_t - t \frac{d\gamma}{dt}\right) (Mv)^{\frac{2}{3}} = K'_t \qquad . \tag{I}$$

and

$$\left(\gamma_t - t \frac{d\gamma}{dt}\right) (x M v)^{\frac{2}{3}} = K_t \qquad . \qquad . \qquad (2)$$

much the same as the equations based on Ramsay and Shields' method. When x = 1, then K' has risen until equal to K. From (1) and (2)

$$x_t = \left(\frac{K}{K'}\right)^{\frac{3}{2}},$$

where K' is the observed value and K the normal, being the sum of the atomic values.

In this way, values of x, calculated from the existing data of other workers, were assigned to the following substances selected as typical examples.

Substance.	T (abs.).	K' (obs.).	K (calcd.).	x.
n-Propylamine	283°-318°	1044-1098	1179	1.30-1.11
Formamide	293°-348°	952-1058	1173	1.32-1.12
Acetamide	363°-385 [.] 5°	1125-1180	1256	1.18-1.10
Acetanilide	398°-425°5°	1830-1850	1936	1.00-1.02
Ethyl urethane	340°5°-370°5°	1240-1280	1416	1.33-1.52

Aliphatic acids and hydroxyl-containing compounds are likewise found to be associated, but owing to the difficulty of deciding on the atomic surface energy to be assigned to hydroxylic oxygen, no values of \boldsymbol{x} were calculated.

Although this method may be admitted as marking a real advance, it raises a number of questions by reason of the difference between its results and those of other methods. Thus, the nitriles appear to be non-associated, the calculated values of K agreeing with the observed, although other methods make them out to be associated. The values of x are also much lower. What is of still greater interest and importance is that the values of x are so much lower than and different in order from the degree of complexity in solution. This question will again be raised in the next chapter.

Methods based on the Specific Cohesion.

Instead of making the surface tension of the liquid the basis of study, the relationship between the surface tension and the density, called the specific cohesion, namely,

$$a^2=\frac{2\gamma}{d},$$

may be used, a^2 standing to represent the specific cohesion.

Several workers on the molecular state of liquids have expressed their measurements in this unit and, curiously enough,

three of them have independently deduced identical rules for the determination of molecular weight involving the specific cohesion.

The specific cohesion which exists in the surface layer of a liquid may be supposed to have a relationship to the amount of heat absorbed during evaporation, and it may be argued that the greater the cohesion the greater must be the amount of heat imported to cause molecules to break through the surface layer and pass into the vapour space. Both Walden ¹⁹² and Dutoit and Moijou ¹⁹³ came independently to the conclusion that the latent heat and specific cohesion would probably be proportional to one another and experiment confirmed this reasoning, as indicated by the following selected results:—

WALDEN.		DUTOIT AND MOIJOU.		
Substance.	$\frac{l}{a^2}$.	Substance.	$\frac{l}{a^2}$.	
Stannic chloride	17:35	Pyridine	18.8	
Silicon tetrachloride	17.03	Benzene	18.4	
Methyl iodide	18.13	Anisole	18.1	
Ethyl iodide	17.90	Hexane	18.8	
Chloral	17.91	Dimethylaniline	18.1	
Carbon disulphide	17.89			

The results are closely the same, being closer still when the mean values are taken of the whole of the substances investigated. By Walden the mean value found was 17.9.

All the above substances are normal; for others, known or suspected of association, such values as the following were obtained (Walden):—

Substance.	$\frac{l}{a^{2}}$	Substance.	$\frac{l}{a^2}$.
Water	44 . 1	Formic acid	22.8
Methyl alcohol	51.31	Acetic acid	25.0 8
Ethyl alcohol	45'27	Acetonitrile	29.17
Ethylene glycol	3 0 •86	Propionitrile	25.75
Phenol	27.21	Butyronitrile	23*35
Acetone	24.12	Amylnitrile	18.96

In all cases, unlike Trouton's Constant (p. 52), the values considerably exceed the normal, and the ratio therefore becomes a good test of association. But neither the extent nor the order of association is capable of being satisfactorily determined and the method might have been included in the previous chapter on qualitative tests but for the fact that it has been converted into one for calculating the actual molecular weight. The process adopted by Walden is to combine the above expression with Trouton's Rule. Thus,

$$l = \text{constant} \times a^2 = 17.9a^2$$
 and $\frac{\text{M}l}{\text{T}} = \text{constant} = 20.7 \text{ (p. 52)}.$

Substituting for *l*,

$$\frac{M \text{ i 7.9 } a^2}{T} = 20.7$$
or $\frac{Ma^2}{T} = \text{ i .162.}$

The quantity Ma^2 is termed the *molecular cohesion*, and the numerical value of the constant I·162 was confirmed by tests with a large number of non-associated substances.

Precisely the same expression had already been arrived at by Kistiakowsky three years earlier, and although it was then deduced empirically, this same worker has recently placed it upon a theoretical basis. 194 Dutoit and Moijou also based their method on the relationship discovered by Kistiakowsky.

The practical measurement required in this method is that of surface tension and density at the boiling-point T, and if these are known, the molecular weight may be calculated. In the preceding chapter reference has already been made to the fact that the mode of variation of the temperature co-efficient of molecular cohesion, $\frac{\Delta \text{ M}a^2}{\Delta t}$, with molecular weight, is very similar

to that of Ramsay and Shields' constant. But although the high values of the co-efficient of molecular surface energy make the Ramsay and Shields' method inapplicable to substances of high molecular weight, this disadvantage does not attend the Walden-Kistiakowsky method, as will be seen from the following table in which the molecular weights derived by the aid of the formula are compared with the sum of the atomic weights.

Substance.	M.W.	т.	a2 × 102.	M.W. (obs.).	
Ethylenedichloride	99	356·3°	4.30	98.4	
Benzyl chloride	126.5	451°	4.20	124.6	
Nickel carbonyl	171	316°	1. 88	195	
Chlorobenzene	112.2	405°	4'13	113.8	
Diphenylamine	169	575°	3.72	179	
Diphenyl	182	557°	3*49	185	
Conylene (CH ₃ . C ₇ H ₁₁)	110	404°	4.062	115	
Rutylidene (C ₄ H ₉ . C ₇ H ₁₁)	152	478°	3.209	159	
Acetylene tetrabromide	168	420°	2.001	160	
Hexaethylbenzene	246	571°	2.00	236	
Diethylsebacate	258	580°	2.414	256	

Nearly all the above substances have abnormally high values of the Ramsay and Shields' constant.

The application to associated substances may now be illustrated by the following table in which x, the degree of association, is $\frac{M.W. \text{ (obs.)}}{M.W. \text{ (calcd.)}}$.

Substance.	x.	Substance.	x.
Water	1.98	Formic acid	1.78
Methyl alcohol	2.40	Acetic acid	1.02
Ethyl alcohol	1.80	Acetonitrile	1.40
Ethylene glycol	1.40	Propionitrile	1.20
Phenol	1,13	Acetone	1.27

One point in favour of the method is that comparison of different substances is made at the boiling-point, an approximately corresponding temperature. Yet a knowledge is desirable of the molecular complexity at other temperatures. Both Walden and Dutoit and Moijou have attempted to extend the formula, therefore. Walden achieved his object by considering the variation of specific cohesion with temperature, the relation being linear and represented by $a^2_t = a^2_o (\mathbf{I} - \mathbf{K}_1 t)$ where \mathbf{K}_1 is the temperature co-efficient of specific cohesion. For some lower temperature t, the formula therefore became :—

$$\mathbf{M} = \frac{\mathbf{I} \cdot \mathbf{I6} \ \mathbf{T}_{\sigma} \left(\mathbf{I} - \mathbf{K}_{1} t\right)}{a_{t}^{2} \left(\mathbf{I} - \mathbf{K}_{1} t_{\sigma}\right)},$$

where T_{σ} is the absolute boiling-point, t_{σ} that on the centigrade scale (since the formula for the variation of specific cohesion is based on a^2 at o°), and a^{2t} the specific cohesion at the desired temperature t.

The formula of Dutoit and Moijou is empirical, namely,

$$M = \frac{0.6 \text{ T } (4.8 - \log p)}{a^2},$$

and investigation at different temperatures is carried out by varying the pressure under which the liquid boils. For a large number of liquids known to be non-associated, the formula yields excellent results over a wide range of temperatures, and pressures from I – I500 mm., the greatest deviation with benzene, chlorobenzene, ethylene dibromide, dimethylaniline, carbon tetrachloride and quinoline not exceeding 3 per cent.

Still another plan adopted by Walden 195 to extend the Kistiakowsky-Walden rule, is based on the idea that the melting-point may possibly, like the boiling-point, be a corresponding

temperature. If this is so, the method would afford information of the molecular state of substances, such as salts and metals, the high boiling-points of which prevent the application of the original formula. The question whether the melting-point may be regarded as a corresponding temperature was tested by experiment, the result being very favourable to such a view. In the revised formula,

$$\frac{Ma^2_{\phi}}{T_{\phi}} = constant,$$

 T_{ϕ} and a^2_{ϕ} being the absolute melting-point temperature and a^2_{ϕ} the corresponding specific cohesion, the mean value of the constant was determined as 3.65. By means of the new formula the following values of x, the degree of association at the melting-point, were obtained.

Substance.	x.	Substance.	x.
Water	3.28	Urethane	2.16
Formic acid	3*4	Sulphuric acid	1.73
Acetic acid	3.3	Benzene	1.82
Ethylene glycol	1.74	Naphthalene	1'46
Phenol	1. 68	•	-

Unlike other methods, this one makes bromine, benzene, and naphthalene appear associated, so that its value, other than as a test of association, is very doubtful.

CHAPTER VII.

MOLECULAR COMPLEXITY IN THE LIQUID STATE.

Some Other Methods and a Review.

Longinescu's Method. 196

THIS is a method depending on the application of an empirical relationship found to exist between the absolute boiling-point, the density and the number of atoms present in the compound.

For any pair of normal liquids of boiling-points T and T', respectively, on the absolute scale, densities d and d' at o° , and containing n and n' atoms per molecule,

$$\frac{\mathrm{T}}{\mathrm{T}} = \frac{d}{d'} \sqrt{\frac{n}{n'}}$$
, or $\frac{\mathrm{T}}{d\sqrt{n}} = \frac{\mathrm{T}'}{d'\sqrt{n'}} = \text{constant}$,

and from the examination of a large number of substances, all normal, the value of the constant was deduced as 100. Conversely, if T and d are known, the number of atoms can be deduced from the formula $n = \left(\frac{T}{100 \ d}\right)^2$ and the complexity of the substance therefore deduced.

The two following tables indicate the first the extent of agreement between the calculated (n) and the theoretical number of atoms (n_0) for normal liquids; the second, the number of atoms present in some associated substances.

I. NORMAL LIQUIDS.

		I. NORMAL	LIQUIDS.		
Substance.	no.	n (calcd.).	Substance.	no.	
Acetyl chloride	7	8	Ethyl propionate	17	16
Chlorobenzene	12	. 12	Phosphorus oxychlor	ide 5	
Ethylene dichloride	6	6	Carbon disulphide	3	6
Silicon tetrachloride	5	5	Sulphur dioxide	3	3
Ethyl cinnamate	25	26	Triphenylamine	32	31
			Octyl ether	51	49
	II.	ASSOCIATE	SUBSTANCES.		
Substance.	no.	*	Substance.	no.	n.
Formic acid	5	9	Water	3	14
Acetic acid	8	14	Hydrogen fluoride	2	9
Methyl alcohol	6	19	Hydrogen cyanide	3	18
Ethyl alcohol	9	19	Ammonia	4	14
Propyl alcohol	12	20	Phosphorus	4 .	9
Glycol	IO	17	Sulphur	8	12
Propylamine	13	20	Sodium	I	108
Phenol	13	18	Potassium	I	132
Acetanilide	6	19			
		7	14		

The method has been applied by its author to a large number of substances, and where comparison is possible, there is very fair agreement with the results obtained by other methods. Extraordinary results are obtained in some cases, notably with sodium and potassium, but this much is to be said, that some of the very high results are obtained for inorganic substances whose boiling points in many cases require revision.

Latent Heat Methods.

Of these methods there are quite a number, but of all of them the best that can be said is that they serve to detect association only, although they have been applied actually to quantitative determination. One such method is that of Walden, 195 depending on the application of Trouton's Rule to substances at the melting-point. Previous experiment had indicated that the melting-point might be regarded as marking approximately a corresponding temperature for different substances and the molecular cohesion method was then extended to this temperature. The success of the latter method suggested that Trouton's Rule might likewise be found applicable at the melting-point, and experiments with thirty-four normal substances indicated that

$$\frac{Ml_{\phi}}{T_{\phi}} = constant = 13.5,$$

 l_ϕ and T_ϕ being the latent heat of fusion and absolute melting-point, respectively. The greatest deviation observed was about 10 per cent.

In turn, the molecular weight could be calculated when the value of this constant had been fixed. When applied to known associated substances and to salts, the following results ensued:

Substance.	x.	Substance.	x.
Water	2.57	Silver chloride	2.3
Formamide	1.62	Potassium nitrate	1.7
Ethylene cyanide	4.40	Lead chloride	r·8
Formic acid	1.44	Lead bromide	2.3
Sulphuric acid	1.41	Aluminium bromide	1.8

Probably objections may be urged against this method as against Trouton's Rule in its ordinary form. The occurrence of high and low abnormal values for associated substances tested at the boiling-point does not appear to be experienced at the melting-point. Fair agreement with the results of the molecular

cohesion method is found, but not with those obtained by Ramsay and Shields' method. Thus, by the latter method, Bottomley 174 found the value of x for potassium nitrate as high as 10, and high values were likewise found by Lorenz and Kaufler. 175

Quite a number ¹⁹⁷ of relationships resembling Trouton's Rule have been deduced and applied to the study of molecular complexity, but none of them serves to do more than indicate cases where association probably occurs.

Bingham's Fluidity Method. 198

This method depends on a comparison of the observed and calculated values of the absolute temperatures at which a given compound has a fixed fluidity (reciprocal of viscosity). If the temperature variation of fluidity is known, it is possible to calculate the temperature at which the fluidity has a fixed value, either 200 or 300 being the comparison value used. calculation of the corresponding absolute temperature is carried out by assigning temperature values to the elements or groups of elements, and from the table so constructed, adding up the atomic or group values for the compound required. Thus, in a homologous series, it is possible to calculate the temperature at which the fluidity of each member is 300, and therefore also the difference of temperature for successive members, i.e. the difference for CH₂. It is then possible, as in atomic refractions or volumes, to assign values to the elements, to a double bond, a benzene ring and so on. Such values, for a fluidity of 300, are the following:-

Carbon	- 110.2	iso-Union	-8.2
Hydrogen	67.8	Double bond	131.3
Oxygen	27.1	Benzene ring	164.3

By the addition of these constants, the absolute temperature at which the fluidity of any particular compound reaches 300, is readily obtained. Should the temperature so calculated agree with that observed, the substance is looked on as non-associated; but should the calculated temperature be lower than the experimental, the ratio of the latter to the former is regarded as the measure of the degree of association. The following association factors (x) have been so obtained.

TEMPERATURE (abs.) OF FLUIDITY 300.

Substance.	Calculated.	Observed.	x
Water	162.7	358.5	2'20
Methyl alcohol	188.1	336.9	1.79
Ethyl alcohol	213.2	371.5	1.74
Acetic acid	236•3	407.9	1.73
Methyl formate	236.3	297'5	1.56
Acetic anhydride	309.9	388·1	1.22
Acetone	207.5	289.5	1'23

Traube's Volume Methods.

According to Traube's measurements, a non-associated liquid, when dissolved in water to form a dilute solution, exerts an attractive influence on the water causing a contraction amounting always to 12.2 c.c. for every gram molecule dissolved. substances which Ramsay and Shields' method indicates to be associated, the contraction brought about is always less than On the assumption that the diminution of the contraction is proportional to the degree of association of the substance, it is possible to find the degree of association. Thus, the contraction observed by the solution in water of 60 grams of acetic acid was found to be 5.8 c.c. Hence, the degree of association =

= 2.1. In a similar manner, the degree of association of a large number of substances, organic and inorganic, has been obtained by Traube. 199

The better known "co-volume method" makes use of the results mentioned in the preceding paragraph, together with a further discovery on the change of volume which occurs when a molecule of a substance is formed. According to Kopp, the molecular volume of any liquid may be represented as the sum of the atomic volumes of the atoms. Traube, from experiments in dilute aqueous solution, showed, however, that the molecular volume is greater than the sum of the true atomic volumes by an amount at 15°, of 13.5 c.c., that is to say, the formation of a molecule in dilute aqueous solution is attended by an increase in volume.

Now as it had already been found that a contraction occurs of 12.4 c.c.* when a gram molecule of a substance is dissolved in water, it follows that the homogeneous liquid has a molecular

^{*} The original value of the contraction was 12.2 c.c. Subsequent experiments led to 12'4 c.c.

volume 13.5 + 12.4 = 25.9 c.c. (at 15°) greater than the sum of the atomic volumes of the elements. The molecular volume of the pure liquid may therefore be written:—

 $V_m = n9.9 \text{ C} + p3.1 \text{ H} + q10' + r40'' + \dots 25.9$ where n, p, q, r, and s are the respective numbers of atoms of each kind, 9.9, 3.1, 2.3, and 5.5 are the true atomic volumes of the elements, O' being hydroxylic oxygen, and O'' carbonyl.

More simply, the formula is

$$V_m = \frac{M}{d} = \Sigma' (At. Vol.) + 25.9$$

where d is the density. The equation expresses the fact that the volume occupied by a substance is the sum of the volume filled by the molecules and the space between the molecules (the covolume).

The co-volume, 25.9, is an average number. For a very large number of unassociated substances, the value was found to range from $22-28.^{200}$ A few substances of high molecular weight, and some tertiary compounds, have rather higher values, the co-volume of tri-isobutylamine being 37.1. When a substance is associated, the co-volume found is smaller than 25.9, if the molecular weight chosen for the substance is the normal value. By inserting the true molecular weight, xM, where x is the degree of association, the co-volume is restored to its true molecular value.

Hence the equation becomes

$$\frac{xM}{d} = x \Sigma (At. Vol.) + 25.9,$$

and x can be calculated by measuring the density at 15° and finding Σ (At. Vol.) from the table subjoined:—

Carbon	9.9	Chlorine	13.5
Hydrogen	3.1	Bromine	13.5
Oxygen (OH of COOH)	0.4	Iodine	13.3
Oxygen (OH)	2'3	Nitrogen (N'")	1.2
Oxygen (CO)	5.2	Nitrogen (NO2)	8.2-10.2

By this means Traube 201 arrived at values of x for a large number of liquids of which the following table contains a selection.

Substance.	x.	Substance.	x.
Water	3.06	Formic acid	1.80
Methyl alcohol	1.79	Acetic acid	1.26
Ethyl alcohol	1.67	Propionic acid	1.46
iso-Propyl alcohol	1.66	Propionitrile	1.14
Glycol	1.88	Acetone	1.23
Glycerine	1.83	Phenol	1.43

The method, apart from any other criticism which may be levelled against it, has an obvious drawback in that a prior knowledge of the constitution of the substance is required before its molecular volume can be calculated.

Guye's Method. 202

The existence of associated molecules in a vapour is regarded by Guye as proof of the complexity of the liquid and, on the basis of the law of mass action and the assumption that van der Waals' equation is applicable to a liquid provided no change of molecular condition occurs on condensation, he has deduced a method of measuring the extent of association.

The case treated was water, but this substance may be regarded as only a type of association represented by the formula

$$A_2 \stackrel{\Rightarrow}{\rightleftharpoons} 2A$$
.

It is presumed that this equilibrium exists both in the vapour and in the liquid; for the former, the molecular concentrations are connected thus,

$$C_2 = KC_1^2, \dots$$
 (1)

and for the latter,

$$C'_{2} = K'(C'_{1})^{2}, . . (2)$$

where C_1 and C_2 and C_1 and C_2 are the concentrations of the single and double molecules respectively in the gaseous and liquid states, and K and K' the respective equilibrium constants.

The question arises whether, knowing the quantities contained in the first equation, it is possible to investigate the second. If K may be substituted for K', then the equation becomes

$$C'_{2} = K(C_{1}')^{2}$$
 . . . (3).

The arguments advanced in favour of the possibility of making this substitution are that it is possible to transform a vapour continuously into a liquid and that in the equilibrium $N_2O_4 \geq 2NO_2$, the equilibrium factor remains constant for considerable changes in the specific volume. With this assumption made, let m_1 be the number of single water molecules, m_2 of double, in a unit volume of liquid. Then equation (3) becomes

$$m_2 = K(m_1)^2$$
 . . . (4).

Now suppose this unit volume of water be converted at constant temperature into v c.c. of vapour containing n_1 single and n_2 double molecules.

Then the equation for the equilibrium in the vapour is

$$\frac{n_2}{v} = K \left(\frac{n_1}{v}\right)^2 \cdot \cdot \cdot \cdot (5)$$

since $\frac{n_1}{n_2}$ and $\frac{n_2}{n_1}$ represent the concentrations of the single and double forms.

Instead of dealing with the actual weight of water, suppose the weight of the single molecule be taken as unit, the mass being the same whether it is a molecule of gas or liquid. Then,

$$m_1 + 2m_2 = n_1 + 2n_2 = N$$
 . . . (6) and if the degree of association of the liquid is x , then

$$x = \frac{m_1 + 2m_2}{m_1 + m_2} = \frac{N}{m_1 + m_2} \qquad . \tag{7}$$

Also, the ratio of the weight of the single to that of the double gas molecules may be written y, where

$$m_2 = K (N - 2m_2)^2$$

= $\left(4 NK + I - \frac{\sqrt{I + 8 NK}}{8K}\right)$, and $m_1 = \left(-I + \frac{\sqrt{I + 8 NK}}{4K}\right)$.

These values of m_1 and m_2 are now inserted in equation (7) so The quantities N and K are got thus:as to give x.

$$\frac{n_2}{n_1} = K \frac{n_1}{v} \text{ by transformation of (5),}$$
and from (8),
$$n_1 = \frac{v}{2Ky} \text{ and } n_2 = \frac{v}{4Ky^2}.$$
Finally,
$$x = \frac{2v(1+y) - y^2 - y\sqrt{4v(1+y) + y^2}}{v(1+y) - 2y^2}.$$

The degree of association can be obtained from the known specific volume v, and the degree of dissociation a of the double molecules of vapour, since $y = \frac{a}{1 - a}$

For water the following values of x were calculated:—

Temperature.	a.	v.	x.
8o°	0.013	3,285 c.c.	1.00
100°	0.011	1,582	1.86
120°	0.010	826.6	1.83

The value of v is measured under the pressure exerted by the vapour at the corresponding temperature.

The weakness of the method lies in the assumption that K can be substituted for K' in the equation applying to the liquid; for van der Waals suggested that the equation bearing his name applied to the liquid only when no alteration of molecular complexity occurred during the transformation of state, and with water this is not fulfilled. Possibly also, molecules of greater complexity than double may be present, although the method might be extended to cover this case. The results, however, agree very well with those obtained at the same temperature by Walden (molecular cohesion method) and by Dutoit and Moijou.

The results also lead to the conclusion that whenever association occurs in the state of vapour, if only to the extent of I in 10,000, association will be appreciable in the liquid state. On the other hand, if a liquid is not associated, then no trace of association of the vapour molecules can occur.

Holmes' Method.203

Both the premises and the results of this method are very different from those already discussed. As a matter of fact, the determination of molecular complexity is based, not on the property of a liquid in the pure state, but by the volume changes which it undergoes when mixed with other liquids. In opposition to the views of many chemists, Holmes believes chemical changes to be rare when two liquids are mixed, the evolution and absorption of heat and change of volume all being supposed to be due to the agency of physical forces acting between the molecules.

The alteration of volume accompanying admixture of two liquids should, on the view that only physical forces are in evidence, be greatest in a mixture composed of equal numbers of each kind of molecule. This view can be made the subject of experimental test. The further assumption on which the method is based, is that the extent to which two liquids mix is governed in a large measure by the relative molecular volumes, or, what comes to the same thing, by the radii of these volumes, assumed to be spherical, in which the molecule exerts an influence.

In support of the first view, it was found that the maximum deviation between the calculated and observed volume of the

constituent of the mixture having the larger molecular volume (referred to the gaseous state) occurred either at equal or very simple molecular proportions. Thus, the maximum difference between the calculated and observed volume of alcohol in an aqueous solution is found when an equal number of alcohol and water molecules are present, and closely similar results are obtained when water is mixed with ethyl and propyl alcohols, formic, acetic and propionic acids and acetone. From these results, the conclusion was drawn that all these substances are of similar complexity, so that if the molecules of liquid water are normal in size, all the liquids named are normal; and if associated, the degree of association is the same. Hence follows the method of finding the relative degree of complexity of a liquid, namely, by mixing it with some other liquid capable of unlimited solubility and determining the composition of the liquid for which the greatest deviation occurs between the calculated and observed proportions by volume of the constituent with the greater molecular volume.

Water cannot be used as one constituent in many cases, but some liquid whose behaviour with water is already known may be employed. Thus, because the greatest deviation occurs with ether and alcohol when mixed in the proportion $(C_2H_6)_2O:C_2H_6O$, ether is regarded as of the same degree of complexity as alcohol and water. In turn, maximum deviations occur with mixtures of ether with benzene or chloroform at proportions $2C_6H_6:(C_2H_5)_2O$ and $2\ CHCl_3:(C_2H_5)_2O$ respectively, and benzene and chloroform are held to have double the degree of aggregation of ether.

The degree of complexity being only a relative value, some standard must be selected and ethyl tartrate was found to serve this purpose because this substance (and glycerol) has a lower degree of aggregation relative to water than other liquids tested. The deviations between the calculated and observed volumes of this ester in water are given below:—

Composit	ion.	Deviation.
$C_8H_{14}O_6$. H	I_2O	0.842
21	I_2O	1.181
31	H_2O	1.325
41	$O_{\mathbf{g}}F$	1.379
51	H_2O	1.362
61	H_2O	1.330
81	H ₂ O	1.202
12]	H ₂ O	0.974

83

The occurrence of the maximum deviation when the composition of the solution is given by $C_8H_{14}O_6$. $_4H_2O$ is, on the theory, an indication that relative to ethyl tartrate, the water molecule is represented by $(H_2O)_4$.

By application of the process to other mixtures, the following table was drawn up. The temperature given is that at which the density of the mixture was determined.

Substance.	Temperature.	Degree of Complexity.
Glycerol	15°	$C_3H_8O_3$
Ethyl tartrate	15°	$C_8H_{14}O_6$
Nicotine	20°	$(C_{10}H_{14}N_2)_2$
Formic acid	20°	$(CH_2O_2)_4$
Acetic acid	15°	$(\mathrm{C_2H_4O_2})_4$
Propionic acid	25°	$(\mathrm{C_3H_6O_2})_4$
Methyl alcohol	15.2°	$(CH_4O)_4$
Ethyl alcohol	15°5°	$(C_2H_6O)_4$
n-Propyl alcohol	25°	$(C_3H_8O)_4$
Acetone	20°	$(C_3H_6O)_4$
Ethyl acetate	25°	$(C_4H_8O_2)_4$
Ethyl ether	25°	$(C_4H_{10}O)_4$
Methyl iodide	25°	$(CH_3I)_8$
Chloroform	20°	$(CHCl_3)_8$
Benzene	20°	$(C_6H_6)_8$
n-Hexane	17°	$(C_6H_{14})_8$
Carbon disulphide	15.2°	$(CS_2)_{16}$.

The results are almost entirely in opposition to those derived by methods based on physical theories and laws current in chemistry and have therefore been subjected to criticism. 204 Some of these difficulties will be clear from the discussion in the next section. Here two objections may be mentioned (1) that since the degree of complexity appears in all cases to be a whole number, then it is almost certain that the liquid cannot be a mixture of two or more types of molecule and further that temperature variation can produce no effect on the degree of association, especially since the liquids are not compared in corresponding states; (2) the values of the degree of complexity are deduced from mixtures of liquids, and the degree of complexity appears to be independent of the second constituent of the mixture, a result wholly different from those based on measurements of the osmotic pressure or depression of freezing point, By the latter methods benzene, for example, has not given evidence of being of complex molecular character.

General Review of Methods and Results.

A general review of the many methods shows that there is a similarity between the state of a given substance in solution and in the liquid condition. As in solution, so also pure liquid substances containing the electro-negative groups OH, CONH, COOH, NOH, CN, NH₂ and CO, namely, water, the alcohols, phenols, amides and anilides, acids, oximes, nitriles, primary amines, and (certain) ketones, are associated, whilst fused salts, according to most methods, are still more associated.

Further, the degree of complexity is never represented by a whole number, so that, as in gases and solutions of associated substances, there must be at least two differently sized molecules. Moreover this degree of association varies with the temperature, decreasing as the temperature rises until, apparently, association disappears. From this circumstance, one may expect that heat is evolved by the association of liquid as of gaseous molecules.

All the above deductions can be made with much confidence; but which, if any, of the methods lead to the actual values of the degree of complexity is quite another matter. The extent of agreement may be tested by reference to substances, such as water, the alcohols and acids, which have been the subjects of tests by so many investigators, and the following table contains data for the comparison, the initials standing for the names of investigators previously mentioned in the text as devising the method.

DEGREE	OF	COMPLEXITY.

Substance.	RS.	RR.I.	в.	вм.	W ₁ .	W ₂ .	DM.	G.	L.
Water Methyl alcohol Ethyl alcohol Formic acid Acetic acid n-Propylamine	2.66 3.24 2.43 3.13 2.9 1.24	1.405 2.17 1.46 — 1.49	1·10 3·43 3·65 2·68 2·74		1.98 2.40 1.80 1.78 1.95	2·57 — — — —	1.96 	1.82	4.6 3.2 2.1 1.8 1.7

The symbol W.₁ represents Walden's molecular cohesion method applied at the boiling-point; W.₂, the same worker's application of Trouton's Rule at the melting-point. It is not always possible to compare the values at the same temperature, but except that under W.₂, all the results for water and the

alcohols are for the boiling-point temperature. Adopting water as the test substance it will be seen that the methods W.₁, D.-M., and G. are in very fair agreement, but these apart, the degree of complexity varies between wide limits. For methyl alcohol, there is fair agreement between R.-S. and B., and between R.-R.I. and W.₁; for ethyl alcohol between R.-R.I. and D.-M. and W.₁ and L. It is therefore obvious that no great importance can be placed on the agreement between two or more methods for any one substance, as the same methods usually lead to different results when tested by a second substance.

In the absence of any guiding principle all that one can hope to do is to select the method against which fewest objections can be raised, and even here opinion is divided. Some methods must clearly be regarded as approximations, such as those depending on the use of melting-point as a corresponding temperature.

Of the surface tension methods the most satisfactory appear to be those of Walden and of Dutoit-Moijou based on specific cohesion. Bennett and Mitchell's method is still sounder in theory, but its results are rather difficult to explain in the light of solution measurements. But against surface tension methods in any form several objections have been raised, notably by Guye, 205 in whose opinion the molecular weight in the surface film is only the same as in the interior when the liquid is normal and not of high molecular weight; whilst if the liquid is associated, the degree of association will be greater in the liquid film owing to the powerful forces acting there. The methods most acceptable are, according to Guye, those which treat the liquid as a whole, like that of Traube or Longinescu. On the other hand, Tyrer 177 would reject surface tension methods, because they lead to results, in his opinion, too low; and, because they do not conform to the following conditions which any method must fulfil, he would also discard Ramsay and Shields', Ramsay-Rose-Innes', Batschinski's, Longinescu's, and Traube's methods:-

- (I) The method must be based on a formula which contains a constant, the same for all substances, and be independent of the constitution.
- (2) The formula must contain the molecular weight as one term.
 - (3) All the terms must be experimentally determinable and

must be related to the same molecular condition, or, if not capable of experimental determination, must be capable of deduction through differentiation or other suitable process.

- 4. All physical quantities must be independent of the number of associated molecular species existing in the liquid; or they must be strictly proportional to the association factor.
- 5. All the physical terms included in the equation must depend on the liquid as a whole and not on localized properties such as surface tension and vapour pressure; and they must either be independent of the vapour phase, or exist under conditions in which the vapour phase may be neglected.

Ramsay and Shields' method does not fulfil (1), (3), and (5); Batschinski's (3) and (5); Longinescu's (3), (4), and (5); Traube's (1). Only the method of Guye (chap. V.) applied at the critical point satisfies the conditions laid down, and this method is limited to a single temperature.

The objection which Tyrer considers vital is that the methods so far proposed are based on the comparison of the properties of a mixture of molecules X_1 , X_2 , and X_3 with a normal liquid of constitution X. But the only knowledge we possess at present depends on selecting some substance as standard and referring the abnormalities of other bodies to probable difference of molecular size; and whether the substances be gaseous, in solution, or liquid, there is no escape from the method.

There is one objection to present methods, however, different either from Guye's or Tyrer's, arising from a want of correspondence between solution measurements and those made on the liquid, an objection previously raised (p. 69) to accepting Bennett and Mitchell's results. The behaviour of alcohol is only one of many cases. In benzene its molecular weight, assuming the validity of Raoult's laws over the range of concentration tested, rises to seven times the normal value, 206 but no method applied to the pure liquid makes the degree of association to be more than about four. In solution the molecular weight rises with increasing concentration and it would be contrary to the law of mass action for a decrease in the degree of association to occur after a certain stage had been reached. Such a discrepancy naturally raises the question whether the values found for any liquid are multiples of the molecular weight in the ideal or gaseous condition, or only of the normal liquid. The actual value assigned to M in any equation applying to a normal liquid is that in the gaseous state. Ramsay ²⁰⁷ was inclined to think at first that the unit of molecular weight was different in the liquid and gaseous states, but modified this opinion in view of the fact that van der Waals' equation may be applied with considerable success to normal liquids, like pentane, ²⁰⁸ and the deduction made is that no change in the molecular state can therefore occur on transformation from gas to liquid.

On the other hand, some physicists regard the liquid state as essentially different from the gaseous in that when condensation occurs complex molecules are always produced; and doubt as to whether the molecular weight of normal liquids is that of the gas is even raised by recent work of van der Waals.²⁰⁹ Both Garver ²¹⁰ and Drucker 211 urge that the normal liquid is really composed of complex molecules. The normal substances, according to Drucker, are those in which the molecules are of a high degree of complexity and remain almost unaffected by temperature alteration, whilst associated substances consist of molecular complexes much more readily broken down. Almost precisely similar are the views of Schames, 212 according to whom the transition from the gaseous to the liquid, as also from the liquid to the solid, is always accompanied by a sharp change in the degree of molecular aggregation. This decision is arrived at by comparing the energy content of the substance before and after condensation at the same temperature. Both benzene and water, in the liquid state, are associated and at each change of state the degree of aggregation is doubled. Near the point of condensation, the saturated vapour of benzene is calculated (at 80°) to have 16.3 per cent of double molecules whilst the liquid itself corresponds to $(C_6H_6)_2$. Near the freezing-point $(C_6H_6)_4$ molecules begin to appear, the solid consisting of (C₆H₆)₄. Liquid water, again, is supposed to contain $(H_2O)_3$ and $(H_2O)_6$; ice $(H_2O)_6$ and $(H_2O)_{12}$ the more complex molecules in each case increasing in proportion. with fall of temperature. Water is supposed to differ from benzene in that it consists, at every temperature, of a mixture of different molecules, whereas with benzene, a type of the so-called normal substance, a mixture is only obtained in the neighbourhood of the change points.

These views, if true, would undoubtedly clear up some of the difficulties arising from a consideration of the results of the various

determinations of the degree of molecular complexity of liquids, but, at present, they can be said only to have reached the suggestion stage. In view of the great influence which van der Waals' equation and its applications have had and still have on our conceptions of condensation and allied phenomena, it is certain that the newer views mentioned will receive the closest scrutiny before being accepted. Undoubtedly before any real advance can be made in our knowledge of the liquid state, we must have first a clear conception of the phenomenon of condensation, particularly in regard to any molecular change that occurs. To accumulate empirical methods of testing liquids is fruitless.

CHAPTER VIII.

THE MOLECULAR COMPLEXITY OF WATER AND THE THEORY OF DYNAMIC ALLOTROPY.

THE great importance of water as a solvent makes a study of its molecular state of particular interest. Whatever its condition as a liquid, its vapour density is not very far removed from the normal and, until recently, the existence of complex vapour molecules had not been recognized, although the vapour densities previously determined were known to contain slight abnormalities. The explanation of these, Bose ²¹³ showed, was due to a small percentage of double molecules in the saturated vapour; for the vapour density results of Kornatz ²¹⁴ could be accurately expressed by the formula

$$d_t = d_o - \frac{K}{P} \left[\sqrt{1 + \frac{d_t P}{K}} - 1 \right] \text{ (vide footnote, p. 19)}.$$

 d_0 was taken to be 1.2432, the value of the density if all the molecules were double. From the density values, the degree of dissociation (a) of the double molecules at various temperatures and under the pressure of the vapour at the corresponding temperature were:—

Temperature.	P (Vap. Pressure)). α.
o°	4.579 mm.	0.934
50°	92°17 ,,	0.018
100°	760°00 ,,	0.011
150°	3581 ",	0.010
200°	11688 ",	0.013

Evidently the dissociating effect of rise of temperature is counterbalanced by the increasing pressure, so that the molecular state remains practically the same over a wide range.

In the state of solution, this tendency to form complex molecules proceeds distinctly further, as illustrated by the following results in various solvents

Solvent.	Concentration.	M.W. $(H_2O = 18)$.
p-Toluidine 215	0.452 -1.602	28.7-33.3
Phenol 215	0.303 -1.350	19.4-19.6
Bromoform 139	0.0102-0.0106	18.9-23.5
Methyl oxalate 139	0.0825-1.820	20.6-35.9
Veratrol 139	0.0230-5.13	17.6-30.0

The fact that liquid water molecules are complex has already been indicated by the many methods described in the preceding three chapters, but there are still other abnormal properties of water, such as melting-point and latent heat of fusion, thermal expansion, specific heat, optical refraction, ²¹⁶ compressibility, viscosity, dielectric constant, ²¹⁷ colour and coefficient of magnetization, ²¹⁸ which afford additional evidence of its abnormal molecular condition.

All the established methods of measurement lead to non-integral values of the factor of association, indicating that two or more molecular forms exist side by side. Before any means of determining association in liquids had been devised, however, Röntgen ²¹⁹ had suggested that all the abnormal properties of water could, qualitatively, be readily accounted for by supposing liquid water to be a saturated solution of ice in some other form of water. These views have been taken up by Sutherland, ²²⁰ who has endeavoured to prove that steam is represented by H_2O , termed hydrol, ice by $(H_2O)_3$, trihydrol, liquid water being a mixture of tri- and dihydrol, $(H_2O)_2$. From the abnormal specific heat, for example, the percentage of trihydrol in liquid water was calculated to be

Temperature o° 20° 40° 60° 80° 100° 120° 140° 198° $(H_2O)_3$ per cent 37.5 32.1 28.4 25.5 23.4 21.7 20.3 19.1 16.5.

The gradual decrease of trihydrol is believed to have its upper limit at the critical point, when water becomes pure dihydrol.

Steam, as already mentioned, has recently been shown to contain some double molecules. At the other end of the scale the triple molecules supposed to constitute ordinary ice, are also supposed to bestow on it its characteristic hexagonal crystalline form through the combination of the oxygen atoms producing a symmetrical molecule.

When water combines with salts, it is supposed by Sutherland to be present as hydrol, because in LiSO₄ $\rm H_2O$, with a single molecule of water, the 18 grams of water occupy a volume, according to density measurements, of 13·3 c.c., much the same as that found for thirty-one other salts containing water of crystallization, in which the mean value is 13·76 c.c. Hence, the density of hydrol when solid is $\frac{18}{13\cdot76} = 1\cdot31$. As ice has a density of 0.917 at 0° and water is a mixture of di- and trihydrol, the density of pure dihydrol may be calculated from the known density of water.

The degree of complexity of liquid water is supposed, according to the same theory, to be altered not only by temperature variation but also when it acts as a solvent, a positive ion converting tri- into dihydrol, and a negative ion causing a reversal of this change; and the failure of the mixture law for the specific heat and surface tension of aqueous solutions is supposed to be due to this transformation of the molecules of solvent in this way.

In somewhat the same manner, Armstrong 221 has attempted to explain the phenomena of aqueous solutions as due less to the solute than the solvent. Instead of limiting the aggregates to groups of three single molecules, rings or chains of molecules may be formed. Of the various types, H_2O , hydrone, and

HO OH, hydronal, are active forms of water, whilst the complexes H₀O: OH₀, H₀O—OH₀, H₀O—OH₀ and so on, are

plexes H_2O : OH_2 , H_2O — OH_2 , H_2O — OH_2 and so on, are OH_2 OH_2

inactive; and the main effect of the solute, it is assumed, is the inter-conversion of and combination of the substance with the various forms.

Bousfield and Lowry ²²² similarly suggest that at moderate temperatures water is a mixture of ice and water molecules, and near its boiling-point, of steam and water molecules, so that water may be a ternary mixture.

Although ordinary ice is the hexagonal form, experiments on the formation of ice under pressure has led to the discovery of several different forms, five stable forms now being definitely recognized. The new forms are denser than ordinary ice, and Tammann ²²³ regards the differences between the various forms as due to different molecular types.

The Theory of Dynamic Allotropy.

The fact that molecules of different kinds can coexist in any liquid, the proportion of each depending on the temperature and other factors, has paved the way for a new view of allotropic modifications, according to which both forms can be present in a state of equilibrium. This theory has arisen more particularly as a result of the study of liquid sulphur, 224 the abnormal behaviour of which has been demonstrated to be due to a mixture of a mobile, yellow liquid called S,, which predominates in amount, and a viscous brown liquid Su the proportion of which increases with rise of temperature and attains a proportion of 40 per cent at the boiling-point, 445°. forms are further distinguished and rendered separable, and therefore capable of determination, by the fact that S_{λ} is soluble, whilst S_u is insoluble in carbon disulphide; S_u, indeed, corresponds in the quasi-solid form to the well-known amorphous sulphur. Certain reagents have the effect of stimulating the rate at which the equilibrium between the two forms is attained at a given temperature, other reagents retard the rate. Ammonia is such a stimulant, sulphur dioxide a retarder, and by the addition of the latter, the mixture can be cooled and afterwards analysed without the equilibrium set up at the higher temperature being disturbed. Sudden cooling by plunging into cold water will also retard the change and preserve the equilibrium proportions. By such a method, Smith and Carson 224 obtained the following proportions of the two varieties:-

Temperature . .
$$121^{\circ}$$
 154° 156° 162° 165° 167° S_{μ} per cent . . 3.75 7.5 8.0 13.5 15.5 16.7

When monoclinic sulphur melts, therefore, it gives rise to two liquid forms, so that at the melting-point (solid) monoclinic sulphur, the two liquid forms S_{λ} and S_{μ} and vapour are all present in equilibrium. The temperature at which the first three are present together under atmospheric pressure is termed the natural freezing-point of monoclinic sulphur, and its value, according to Smith and Carson, ²²⁵ is 114·25°, S_{μ} to the extent of 3.6 per cent being present. Monoclinic sulphur, which is regarded as the solidified S_{λ} , has, when pure, a higher freezing-point, 119·25° according to Smith

and Carson, or 118.95° according to Wigand. The freezingpoint was obtained by the analysis of samples of sulphur which had been heated for varying periods, thereby lowering the freezing-point according to the amount of Su produced, and extrapolating the temperature of freezing to the point at which $S_{\mu} = 0$. When this freezing-point of pure monoclinic sulphur had been fixed, it was possible to calculate the molecular weight of S, from the known depression of freezing-point produced by known quantities present. The atomic depression, that is the lowering of freezing-point by the solution of 32 grams of S_u in 100 grams of S_{λ} , was thus found to be 46.1. Using Person's value of the latent heat of fusion, l, 9.368, the atomic depression calculated from van't Hoff's equation $C = \frac{0.0198 \text{ T}^2}{7}$ was 325°, which is 7.6 times the observed value. Hence Smith and Holmes concluded that the molecular weight of S,, like that of S_{λ} , corresponds to S_{8} , and that the change which accompanies the conversion of one form into the other is intramolecular.

As the value of the molecular size depends on the latent heat of fusion employed, Smith's result has been subjected to revision. Thus Tammann found l=12.5 and the number of atoms 5.74, or, S_{μ} is S_{6} ; Wigand ²²⁴ found l=10.4, and the number of atoms 6.9, and considering this value somewhat low, concluded that S_{μ} is S_{8} . Smith and Carson ²²⁵ using Lewis and Randall's ²²⁶ value for the latent heat of fusion, subsequently identified S_{μ} with S_{6} , as also did Beckmann ^{226a} independently.

This work on the nature of liquid sulphur, coupled with the previously stated views on water, another substance which must be regarded as exhibiting this dynamic allotropy, may have considerable influence on the subject of molecular association along two lines. In the first place it may be possible, as with water and sulphur, to trace the change of molecular condition, if any, through from the gaseous to the solid state. Although it had no stated reference to the theory of dynamic allotropy, Guye's method marks one such attempt since the degree of association of a liquid is determined from the known association in the vapour state. A somewhat different example, and one more clearly illustrating the point, is Preuner and Schupp's ¹⁰ investigation of liquid sulphur. These investigators believed that S_{μ} has the molecular formula S_6 and is identical with the vapour

molecule S_6 already shown to exist in conjunction with S_8 at temperatures near and below the boiling-point. In order to test this view, they calculated from the equilibrium formulæ given in chap. II. the proportion of S_6 present at different temperatures and compared the results with the amounts of S_μ found by Gals. Fairly good agreement was found between the values of Gals and those calculated from the partial pressures of S_6 in the vapour. Thus:—

Temperature 215° 230° 245° 260° 290° 320° Per cent S_{μ} (Gals.) 18 22 25 28 34 37 Per cent S_{6} (P. and S.) 20 22 24 25 30 34

A further attempt, in this case, to connect the liquid and the solid state, has been made by Tammann, 228 based on the coefficient of surface energy of the liquid substance. If a vapour contains molecules of different kinds, then the liquid formed on condensation should be associated and a mixture of two or more molecular species; so also, when the liquid is converted into a solid, the different liquid molecules may be expected to give rise, as a rule, to different crystalline forms. In the main, Tammann finds this is true, that normal liquids only give rise to a single crystal group, whilst the associated give rise in most cases to two stable groups. The test of association is the value of the coefficient of molecular surface energy, association being presumed if k < 2.00. Thus:—

Substance.	No. of crystal groups.	No. of unstable forms.	k.
Water	2	2	o·88
Acetic acid	2		0.00
Formic acid	r	4	0.00
Sulphur	2		1.21
Phenol	2		1.80
Nitrobenzene	r		2.53
p-Cresol	I		2.24
Acetophenone	r		2'40
Diphenylamine	I		2.62
Veratrol	r		2.97

In this way we have the different molecular species preserving their individuality through changes of state, the only effect being an alteration of the proportions of each.

The idea of dynamic allotropy, as with liquid sulphur, has been extended to liquid phosphorus, ²²⁹ and attempts, less successful, have been made to apply the theory even to the solid form of the element. ²³⁰ Benedicks, ²³¹ again, sees in the fact that the

transition point of silver iodide varies over a range of temperature in the neighbourhood of 147°, evidence of the co-existence of two solid forms, mutually soluble and in equilibrium; and he has extended the same mode of explanation to the allotropic forms of iron, reviving the earlier theory of Osmond and Werth that iron exists in two forms only, the present a and γ forms, and that β iron is only a solid solution of these two forms. Further, it has been suggested that the existence of different forms of tin may be due to a similar cause, as also the existence of two phases, the hard and soft states, which many metals can possess. It is well to bear in mind, however, that these are not cases which can be explained on the basis of the existence of dynamic allotropy in the liquid state, for as already shown in preceding chapters, the liquid metals appear to be monatomic.

The second possible effect of the theory on the study of molecular association lies in the investigation of the effect on the solvent of added substances. The experiments on liquid sulphur have brought to light the fact that certain reagents, such as ammonia, may assist in establishing equilibrium between the different molecular varieties; others, like sulphur dioxide, may retard; whilst others still, like iodine, may even displace the point of balance. If this is so with sulphur, there seems no reason why it should not occur with water or other complex liquids when solutes are added, in some such way as already suggested by Sutherland, Armstrong, Bousfield and Lowry, Drucker and Kassel, and the writer (see chap. IV.), all from independent points of view. Very few direct tests of such a possibility have been made.

CHAPTER IX.

THE SELECTION AND USE OF MOLECULAR FORMULÆ.

In order to ascribe a formula to a substance, its composition and the atomic weights of the constituent elements must be known. When known, the simplest possible formula can be calculated and the molecular formula can be obtained by comparing the simplest formula weight with the molecular weight derived by applying Avogadro's hypothesis.

Avogadro's hypothesis requires the comparison of the standard gas and the gas of unknown molecular weight to be carried out under the same conditions of temperature and pressure. As no two gases alter in volume in precisely the same way when heated or compressed, the comparison is carried out either at zero pressure or at a high temperature, and the relative densities then reduced to 0° and 760 mm. (and latitude 45°); or, what comes to the same thing, the deviations from the gas laws between zero pressure and 760 mm., or between 0° and the comparison temperature may be determined and a correction introduced.

The number of accurate molecular weights that can be so deduced is very small; but the molecular formula of a substance can be obtained from an approximate molecular weight determination. Although this approximate method provides molecular formulæ for many substances, there are two classes which create special difficulty, (I) those which cannot be vaporized, (2) those whose molecular size varies with the temperature and pressure.

The first class comprises a very large number of substances, in fact, the majority of known substances. When formulæ such as NaOH, Na₂O, KNO₃, CaSO₄, etc., as well as those of many organic substances, are written down it must be remembered that they may not, in fact, very likely do not, represent molecular quantities, but are only empirical. The true formulæ are (NaOH)_n, (KNO₃)_n and so on, where n is yet unknown. Struc-

tural formulæ both in inorganic and organic chemistry are written for many substances the molecular weights of which are not known, and although much help in studying chemical reactions is often to be gained from their use, it has to be borne in mind that they can only be tentative so long as the molecular formula remains unknown.

The second class of substances illustrates the fact, clearly brought out in the preceding pages, that the molecular weight of a substance is to be regarded as a physical property, dependent, like other properties, on the conditions of temperature, pressure and surroundings. Even for the permanent gases, comparison at 0° and 760 mm. is not easy because of the difficulty of obtaining "the same conditions of temperature and pressure"; and with substances which change their molecular size according to the conditions imposed, the selection of a single formula is obviously impossible. In such cases, the molecular formula selected has usually been the simplest one, on the ground that rise of temperature tends to produce the single molecule. Thus, Biltz and Meyer, 38 when determining the molecular weight of stannous chloride, found the density changed from 8.76 to 7.08 (relative to air) between 639° and 1113°. As the value corresponding to SnCl₂ was 6.53, the investigators named came to the conclusion "dass Zinnchlorürdampf mit Molekülen von der Formel Sn₂Cl₄ in der That nicht existiert". The obvious conclusion is rather that a mixture of molecules SnCl2 and Sn2Cl4 exists over the above range of temperature.

The habit of adopting the limiting value of the formula, or that which seems most likely at high temperatures does not, however, appear to have been very consistently followed. Thus, although stannous chloride, aluminium chloride, and acetic acid are usually written as $\mathrm{SnCl_2}$, $\mathrm{AlCl_3}$, and $\mathrm{CH_3COOH}$ respectively, arsenious oxide and iodine are most often written $\mathrm{As_4O_6}$ and $\mathrm{I_2}$, although in both cases rise of temperature ultimately brings about complete dissociation. Vapour density determinations carried out in the last few years make it clear that most molecular formulæ for elements should, if chosen as above, correspond to the atom. Thus, iodine, bromine, and (apparently) chlorine, sulphur, selenium, hydrogen may all be represented by monatomic molecular formulæ I, Br, Cl, S, Se, and H, and it is most probably only a matter of overcoming experimental difficulties before all elements are reduced to the monatomic state,

The extension of the gas laws to solutions provides a means of testing a large number of the substances belonging to the first class above, namely, the unvaporizable substances, but greater care needs to be taken in interpreting the results than has been observed in the past. In the state of vapour, the factors which may modify the density are the temperature and the pressure. In Meyer's method of finding the density a filling gas is employed, but it is always chosen so as to be chemically indifferent to the vaporized substance. Further, the dielectric constant of a gas or vapour is only so very little greater than unity (vacuum = I) as to be without effect in modifying the forces acting within the molecule. In a solution, the modifying factors are temperature, concentration, and medium, and if vapour density measurements are to be comparable with solution measurements, the following conditions ought to be observed:—

(a) Reduction of the molecular weight to the temperature of o°. (b) The standard solution should be that which exerts an osmotic pressure of one atmosphere. (c) The dielectric constant should be low.

It is obviously not easy to fulfil condition (a), but (b) can more easily be adopted. In addition to (c), care ought to be taken that the solvent shall have no chemical influence on the solute, just as the filling gas in Meyer's method must be an indifferent one. No one would accept vapour density determinations by Meyer's method if oxygen or chlorine was the filling gas. Yet the necessity of this factor is often over-looked in solution measurements, and wrong interpretations of molecular weight results have often been made. Thus, cuprous chloride both in bismuth 233 chloride and in pyridine 120 appears to have the formula CuCl. In bismuth chloride the result is undoubtedly due largely to the high dielectric constant of the solvent; in pyridine, combination with the solvent to form a compound Cu₂Cl₂. 6 C₅H₅N²³⁴ occurs, and the experiment, as a molecular weight determination, ought to be ruled out. The same objection, namely that chemical action occurs, may also be raised to certain of Beckmann's determinations such as those of the molecular weights of a number of metallic salts in quinoline.²³⁵ In the state of vapour, no distinct reduction below the formula Cu₂Cl₂ has so far been obtained, and this formula is probably more truly representative of the molecular state of cuprous chloride than the result of the

determination in bismuth chloride, owing to the modifying effect of the dielectric character of this solvent.

From what has been said, the difficulty of obtaining really representative molecular formulæ will be evident. There are several conceivable ways of deriving such formulæ:—

- I. Abandon solution measurements and rely on vapour density determinations only, as in the original application of Avogadro's hypothesis, but carry out the comparison of densities at a temperature within 100° of the boiling-point, since this is roughly a corresponding state for different liquids.
- 2. Rely only on vapour density determinations and select as molecular formula the value corresponding to the limiting density obtainable at high temperatures.
- 3. Make the molecular formula correspond to the particular conditions under which the substance exists.

Obviously (I) and (2) are only compromises. Nothing has yet been said in this chapter of the molecular formulæ of liquids and solids, but (3) takes these states into account. However difficult of attainment, (3) is the only theoretically sound method. It requires that the formula of a substance shall be recognized as a function of the physical conditions under which it exists, so that the formula for elements is X_n , for compounds $(XY)_n$, where n is to be specified according to the conditions. Although we are far from realizing how n varies for most substances, the adoption of such formulæ would bring home clearly the fact that both the physical and the chemical behaviour of a substance may be closely dependent on the value of n.

CHAPTER X.

MOLECULAR ASSOCIATION AND PHYSICAL PROPERTIES.

As shown in preceding chapters, the marked effect which association has on the physical properties of a substance has been employed to provide methods both of detecting and measuring the extent of association. Reference has been made to the abnormal osmotic pressure, lowering of vapour pressure and freezing point, latent heat of evaporation and fusion, thermal expansion and specific heat, critical temperature and the critical ratios, surface tension and surface energy, and viscosity and fluidity. There are still other properties which association has been found to affect, although to an extent which is not sufficiently well marked as to serve, in turn, as a means of detecting association. Several such properties are discussed in this chapter.

Degree of Association and Dielectric Constant of a Liquid.

On the whole, these two properties run parallel, substances strongly associated having, with few exceptions, high dielectric constants. The converse of this is not true, for a number of substances of high dielectric constant are unassociated. These facts are brought out in the following table.^{235a} In the absence of any reliable method widely applied in the determination of degree of association, the association factors obtained by use of the Ramsay and Shields' method are inserted. Too great a value must not be attached to them.

Substance.	Association Factor.	Dielectric Constant.
Formamide	6.18	>84
Ethylene cyanide	>7'3	61.3
Water	3.81	81.7
Formic acid	3.01	58
Acetic acid	3.62	6.3
Methyl alcohol	3'43	32.5
Ethylene glycol	2.02	41'2
Ethyl alcohol	2.74	26°0
Acetamide	2.47	59'2
Propyl alcohol	2*25	22.3

PHYSICAL PROPERTIES

Substance.	Association Factor.	Dielectric Constant.
Lactonitrile	2.21	3 7° 7
iso-Amyl alcohol	r.92	10.0
Acetonitrile	1.73	38.8
Formanilide	1.69	20.5
Nitromethane	>1.20	38.2
Propionitrile	r•48	27.5
Butyronitrile	1.38	20.3
Dimethylnitrosoamine	1.34	53*3
Acetone	1.36	21.2
Acetanilide	1.31	19.5
Acetyl chloride	1.06	15.2
Ethyl thiocyanate	1.04	19.2
Benzylnitrile	1.03	18.3
Phosphorus trichloride	1'02	4'7
Carbon tetrachloride	${f r}$ or ${\cal U}$	2°25
Ethyl iodide	I.oI	7.4
Benzene	I.oI	2.3
Acetic anhydride	0.99	20.7
Ether	0.00	4*3
Benzaldehyde	0°97	14.2
Sulphur monochloride	0.92	4.8
Nitrobenzene	0.93	35*5

It may be remarked that Dutoit and Moijou find nitrobenzene and benzonitrile slightly associated.

Solvent Power and Degree of Association.

Belief in the dependence of solubility on the molecular state has been expressed by a number of observers. One of the first was Crompton, who pointed out that unimolecular liquids mixed readily, but the mutual solubility of unimolecular and associated substances is, as a rule, limited. Walden also found that the solubility of a salt, tetraethylammonium iodide, was distinctly dependent on the degree of association of the solvent. In the following table the degree of association is again that derived from Ramsay and Shields' method and the solubility is the amount of salt per 100 c.c. of solvent.

Solubility at 25°.	Degree of Association.
35*5	3.6
10.4	3'43
7.55	2.02
5*94	1.20
3.04	1.73
o•88	2.74
0.81	1.48
ate 0.475	1.19
0.467	1.02
0'249	1.22
0.0004	0.99
	35.5 10.7 7.55 5.94 3.04 0.88 0.81 ate 0.475 0.467 0.249

Associated substances like formamide, formic and acetic acids, and phenol also readily dissolve this and similar salts, whilst non-associated liquids as ether, benzene, ethylene dibromide, carbon disulphide do not.

Crompton's views have been extended by the writer,²³⁸ who has indicated that a substance is associated, as a rule, only in the solvents in which its solubility is rather limited. Thus, speaking broadly, hydroxyl-containing substances, amides and their derivatives, oximes, nitriles and salts are but slightly soluble in the hydrocarbons and their halogen derivatives, in which association is exhibited. An attempt to obtain a quantitative relationship, however, has not so far succeeded.²³⁹

The belief in the relationship between degree of association and solubility has been carried much farther by Holmes, so far, indeed, as to base a method of determining relative association factors on the extent of miscibility (see chap. VII.)

Electrical Conductivity and Degree of Association.

Under this heading may be included both the power of forming a conducting medium and of acting as an electrolyte in solution.

Associated liquids when acting as solvents usually form good conducting media, and also possess a distinct or high conductivity when pure. Organic liquids certainly cannot be compared with inorganic in this respect, but of the five best conducting liquids found by Walden,²⁴⁰ namely,

Substance. Formamide. Acetamide. Dimethylnitrosoamine. Acetylacetone. Formic Acid. Sp. Conductivity 4.7×10^{-5} 29×10^{-5} 3×10^{-5} 1.6×10^{-5} 1.5×10^{-5} only acetylacetone is non-associated. The conductivity of inorganic halogen-containing compounds is likewise great when the degree of association is highest and drops to zero with non-associated substances. In the following table, the substances are arranged in the order of their degree of association. K represents the specific conductivity. 241.

Substance.	κ.	Substance.	к.
Sodium chloride	3°34 (750°)	Sulphuryl chloride	0.18 × 10-6
Potassium chloride	1.91 (250°)	Arsenic trichloride	1.24 × 10-6
Lead chloride	2°39 (520°)	Tin tetrachloride	Nil.
Silver chloride	1.83 (500°)	Antimony pentachloride	Nil.
Antimony chloride	$109 \times 10^{-6} (80^{\circ})$	Silicon tetrachloride	Nil.
Phosphorus oxychloride	2.2 × 10-6	Phosphorus trichloride	Nil.
Thionyl chloride	2'0 × 10-6		

The conductivity of an electrolyte also depends to a considerable extent on the degree of association of the solvent.²⁴⁰ This follows from the connexion already shown to exist between association factor and dielectric constant, for on the latter property the conductivity of a solution closely depends. As there are many exceptions to the parallelism between association factor and dielectric constant, so likewise the association factor of the solvent does not control the conductivity of the solution.

Inorganic electrolytes are, as a class, associated substances. This statement applies to salts and acids * for, as yet, nothing is known of the molecular condition of bases. Even organic electrolytes are most often associated bodies. Thus, when dissolved in the liquefied halogen haloids, organic acids, alcohols, phenols, amides, and nitriles conduct a current, although ether and thymol, unassociated substances, behave similarly. Although but qualitative, the connexion between association and conductivity and association and ionizing power seems so certain, that of about one hundred associated substances of all types, all have been found to possess a distinct conductivity, or to exert ionizing power, or both. To

In solvents of low dielectric constant, the highly complex molecular state of salts must be taken into account in attempting to explain the conductivity of the solution. An interesting example, in this connexion, is furnished by Isbekow's work ⁸⁷ on aluminium bromide as a solvent, in which non-conducting haloid bodies, such as phosphorus trichloride, were found to be non-associated; whilst salts, like bismuth tribromide, producing conducting solutions, were strongly associated. The molecular conductivity of the salt solutions increased as the concentration increased, so that the ions cannot be of the same type as those which are assumed to carry the current in aqueous solution.

Molecular Association and Optical Rotation.

The effect of the solvent on the rotation of an optically active substance has been the subject of repeated investigation. As the outcome of an extended review of the subject, Walden ²⁴³ came to the conclusions that (1) There exists a connexion between the osmotically determined molecular size of the active solute and its

^{*} The term acid refers here to the dry substance.

rotation in the same solvent, (2) The relationship between the degree of association and the rotation is not, however, revealed in a simple parallelism, but each constituent of the solution exercises constitutive influences on the value of the rotation.

The substances used by Walden in the test were dimethyltartrate, diethyltartrate, dimethylmalate, diethylmalate, and dimethylacetylmalate. With dimethyltartrate the following results were obtained:—

Solvent.	Benzene.	Et. Acetate.	Acetone.	Me. Alcohol.	Acetonitrile.	Chloroform.
M.W.	305	219	195	183	173	253
$[\alpha]_{D}$	- 1.7	+ 2.6	+ 5.8	+ 6.7	+ 10.8	- 6.3

The molecular weight calculated from the simple formula is 178·I, and it will be seen that as the observed molecular weight increases, the rotation falls. As chloroform does not fall into line, and as, moreover, a substance may cause different rotations in solvents in which its molecular weight is normal, it is evident that association cannot be the factor controlling the rotation.²⁴⁴

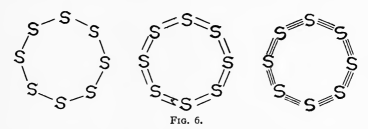
Other evidence of the influence of molecular size on optical activity has been obtained by Navassart ²⁴⁵ who found that the rotation of tannic acid in different solvents appears to decrease as the molecular size of the solute increases.

CHAPTER XI.

MOLECULAR ASSOCIATION AND CHEMICAL COMBINATION.

IN seeking to trace the nature of the forces which cause the molecules of so many substances to form aggregates, one naturally contrasts association with chemical combination. In both phenomena there is attraction between molecules, in the first attraction between like, in the second between unlike molecules. This apparent similarity has, indeed, led several chemists to propound "association" theories of chemical action. When two bodies react, according to this view, at least two stages are passed through. There is first of all a union (or association) of the two molecules resulting in a molecular compound which is not stable, so that a rearrangement of the atoms proceeds with the production either of a new type of molecule or of two or more molecules. Kekulé 246 was the propounder of such a view. H. E. Armstrong ²⁴⁷ explained the oxidation of hydrocarbons in a similar manner whilst Engler 248 suggests that the first stage of all oxidations by oxygen is the association of the oxygen molecule as a whole with the substance being oxidized. solution of ammonia in water is also supposed to result first in the formation of NH3.xH2O, part of which is then transformed into the hydroxide NH₄OH.²⁴⁹ The test of such a hypothesis as Guye 250 pointed out, can readily be made in connexion with many reactions between organic bodies, by investigating their freezing points when mixed, since at such temperatures the reaction is usually exceedingly slow. By a study of their equilibrium diagrams, Wroczynski and Guye 2500 and Schmidlin and Lang 250b have found direct confirmation of this view of the mode of chemical action between a number of organic substances.

Belief in the similarity of molecular association and chemical action also finds expression in the structural formulæ which have been proposed for associations or aggregations of similar molecules. Water and hydroxyl-containing molecules are supposed to form aggregates through oxygen acting as connecting link (see chap. VIII.); whilst for the sulphur molecule, one of the three following formulæ is supposed to represent it ^{250c,} according as the sulphur atom is assumed to be di-, tetra-, or hexavalent.



The best way of illustrating the relation between association and chemical combination is, perhaps, to consider the properties of the components from which molecular compounds are formed. For this purpose the following list, containing, in the main, substances which are associated, has been drawn up. Each pair is only a type of many similar ones.

Constituents.

Hydrogen iodide: Water ²⁵¹
Nitric acid: Water ²⁵²
Potassium hydroxide: Water ²⁵³
Ammonia: Water ²⁵⁴
Lithium iodide: Water ²⁵⁵
Lithium iodide: Me. alcohol ²⁵⁶
Lithium iodide: Et. alcohol ²⁵⁶
Lithium iodide: Pr. alcohol ²⁵⁶
Lithium iodide: Pr. alcohol ²⁵⁶
Lithium iodide: Pr. alcohol ²⁵⁷
Lithium iodide: Acetic acid ²⁵⁷
Lithium iodide: Acetic acid ²⁵⁸
Hydrogen chloride: Formamide ²⁵⁸
Sodium iodide: Acetamide ²⁵⁹
Lithium chloride: Ethylamine ²⁶⁰

Magnesium iodide: Et. carbamate ²⁶¹
Magnesium bromide: Urea ²⁶¹
Magnesium iodide: Acetonitrile ²⁶²
Magnesium bromide: Acetone ²⁶²
Magnesium Iodide: Ammonia ²⁶³
Phenol: Urea ²⁶⁴
p-Cresol: Aniline ²⁶⁴
Phenol: a-Naphthylamine ²⁶⁴
Resorcinol: p-Toluidine ²⁶⁵

Catechol: a-Naphthylamine 265

Compounds Formed.

HI.3H2O; HI.2H2O HNO3.2H2O; HNO3.H2O. KHO. H2O; KHO. 2H2O; KHO. 4H2O $NH_3.H_2O$; $2NH_3.H_2O$ LiI.3H₂O; Li, H₂O. Lil.3CH₃OH LiI.4C2H5OH LiI.4C3H7OH LiI.4C5H11OH 2HBr.C₃H₇OH LiI.3CH3COOH 3HC1: H.CONH2. NaI.2CH3.CONH2 LiCl. $C_2H_5NH_2$; LiCl. $2C_2H_5NH_2$; Li Cl. 3C2H5NH2 $MgI_2.6NH_2.COOC_2H_5$ $M_gBr_2.6CO(NH_2)_2.$ MgI2.6CH3CN $M_gBr_2 \cdot 3(CH_3)_2CO$. $MgI_2.6NH_3.$ $C_6H_5OH.2CO(NH_2)_2$. $C_6H_4CH_3OH \cdot C_6H_5NH_2$. C6H5OH.C10H7NH2. $C_6H_4(OH)_2$. $C_6H_4CH_3NH_2$ $C_6H_4(OH)_2 \cdot C_{10}H_7NH_2$.

Constituents.

Aniline: Et. acetate 248

Phenol: Acetone ²⁴⁹ Resorcinol: Acetone ²⁴⁹

Phenol: Cyclohexanone 249

Urea: p-Nitrophenol 267

l- and d-Dimethyltartrates 266

Compounds Formed.

 $\begin{array}{c} {\rm C_6H_5NH_2\cdot CH_3COOC_2H_5}; \ \ {\rm _3C_6H_5NH_2\cdot} \\ {\rm _2CH_3\cdot COOC_2H_5}; \ \ {\rm _2C_6\cdot H_5\cdot NH_2\cdot CH_3} \end{array}$

COOC₂H₅

 ${}_{2}C_{6}H_{5}OH \cdot (CH_{3})_{2}CO \cdot C_{6}H_{4}(OH)_{2} \cdot 2(CH_{3})_{2}CO \cdot C_{6}H_{5}OH \cdot C_{6}H_{8}O \cdot$

 $C_6H_{10}O_6$. $C_6H_{10}O_6$.

 $CO(NH_2)_2 \cdot C_6H_4NO_2 \cdot OH$.

Ethyl acetate and a-naphthylamine are the only substances which are doubtfully associated; magnesium bromide and iodide have not been tested but are probably associated.

Hundreds of molecular compounds formed from constituents like those above are known and therefore serve to show that associated substances are to a marked degree chemically active. Further, just as with molecular association, so also the formation of additive compounds occurs least readily with ortho-substituted benzene derivatives and much more readily with meta- and paraderivatives. Again, the organic oxygen compounds which form the most stable compounds with salts are those in which the oxygen is attached to hydrogen, whilst, in general, the organic substances either of high dielectric constant or high association factor form the most stable addition compounds with magnesium bromide or iodide. Menschutkin 269 found the order of decreasing stability of these addition compounds as below:—

Urea Ethyl alcohol Ethyl orthoformate Water Acetonitrile Benzaldehyde Aniline Acetic anhydride Methyl acetate Acetamide Acetic acid Acetal Ethyl urethane Propionic acid Acetyl chloride Formic acid Ethyl acetate Ether Methyl alcohol Acetone Anisol

Although it is clear that associated substances are so often chemically active, it is not necessary, however, that either or both constituents of an additive compound shall be associated. In the above list, the substances in the last column are unassociated, and the order of stability in the other two columns is not the order of the degree of association. More convincing is the fact that many additive compounds are formed from unassociated constituents. Among organic substances, examples of this type are the compounds of trinitrobenzene with the orthometa- and para-toluidines, the benzidines, xylidenes and tolidenes; 270 of napthalene, phenanthrene and fluorene with di- and

trinitrobenzenes and trinitrotoluene; ²⁷¹ of ethyl bromide with bromine, ²⁴⁸ benzyl chloride with methylaniline, ²⁴⁸ ethyl acetate with bromine ²⁷² and many others.

Then, on the other hand, substances known to be associated, such as formamide and water, may be mixed without combination occurring.273 It has usually been supposed that what happens in such cases is a dissociative action, that the forces which cause the molecules of any pure substance to form aggregates exert such an attraction on a different aggregated molecule as to cause disruption of each. It has been pointed out in an earlier chapter (p. 48) that such an idea is not generally true. Some very surprising results obtained by English and the writer 274 indicate how very wide of the mark such an idea may be, since, instead of bringing about dissociation, an increase in the degree of association results, at any rate when salts are mixed. Thus, triethylammonium chloride and triethylammonium bromide are each strongly associated in bromoform. When introduced together into the same solvent, the depression of freezing-point observed ought to be greater than the value calculated from the independent depressions if mutual dissociation occurred. Instead, the observed was always less than the calculated depression by an amount which reached a maximum at a certain concentration and then declined, as will be seen from the following table. Δ_a and Δ_c represent the observed and calculated depressions. concentrations of the substances are expressed as grams per 100 of solvent, the chloride being dissolved first and then successive additions of bromide made.

		Concn.	\mathbf{D}_{o} .	$D_c - D_o$.
r.	Chloride	1.693	0.775	_
	Bromide	0.3601	0.813	0.118
		o•6368	o•836	0'224
		1.625	0.958	0.315
2.	Chloride	1.728	o•786	_
	Bromide	2.261	1.163	0.275
		3.469	1.362	0.224
		5.044	1.818	0.010

The extent of the deviation depends on the solvent. In water it is considerably less, but no enhanced dissociation is found to occur.

Evidently associated molecules are centres of very considerable activity. In some cases chemical combination occurs when two associated substances are brought together, in others

(salts in particular) there appears to be repulsion rather than attraction between the molecules. Unsaturation, as measured by the power of chemical action, cannot satisfactorily be accounted the cause of association. The mere presence of trivalent nitrogen, or of elements like chlorine, bromine and iodine, oxygen and sulphur does not bring about molecular association. It is only when these elements are present in one of the electronegative groups mentioned in chaps. III. and VII. that association occurs. Theories of valency which have up to the present been applied to explain the occurrence of molecular association are all unsatisfactory because they are founded on the supposed behaviour of unsaturated elements.⁷⁷

A study of the factors which influence both molecular association and molecular compound formation also reveals some similarities and some differences between the two phenomena. effect of rise of temperature is in both cases to cause increased dissociation. Pressure or concentration likewise affects each similarly; thus, the degree of combination between phenol and α-naphthylamine in benzene solution 274 increases as the concentration becomes greater, whilst, as already shown (chap. III.) increase of concentration raises the degree of molecular aggregation. The effect of the solvent, however, appears to be different in the two phenomena. On molecular association, the dielectric character of the solvent exerts a dominant influence. The influence of the solvent on the combination between triethylamine and ethyl iodide appears to have no reference to its dielectric constant 275

All these facts taken together have led to the suggestion, ⁷⁷, ²⁷⁴ that the forces causing molecular association are either different in origin from those which bring about chemical action; or, if of the same origin, then they act differently. Possibly in molecular association we have to do with electrical as distinct from chemical forces.

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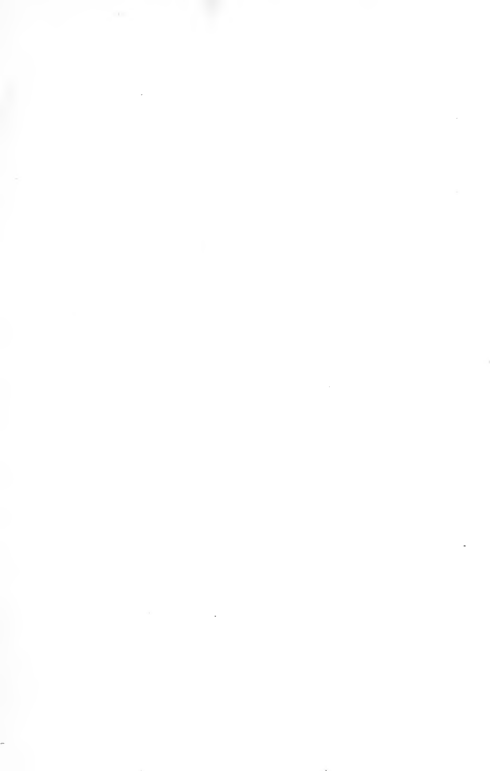
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APPENDIX.

This appendix, as previously mentioned in the text, contains a summary up to date of the various investigations of molecular complexity of dissolved substances, with the concentrations calculated in most cases in terms of grams of substance per 100 grams of solvent. In a few cases the concentration is expressed in terms of milligram molecules per 100 c.c. of solvent (N), or, as milligram molecules per 100 grams of solvent (n). Where the molecular weights were not recorded by the investigator, they have been calculated using the depression or elevation constant quoted in the paper.

Some of the substances quoted have been investigated, in the same solvent, by a number of observers. In such cases all references are given but only one set of figures is recorded, namely, that drawn from the first reference quoted.

The nomenclature adopted is not always that in common use in current English chemical literature, but is, with slight variation only, that given by the investigator of the substance.

It is not easy to define clearly where molecular association exists. The majority of the substances in the table are distinctly associated. Other results are quoted, however, to illustrate the effect either of substitution on the degree of complexity, or of the influence of the solvent. At the end of the appendix a list of substances, or classes of substances, which may be considered as normal, is given.

High molecular weight results are also obtained when solvent and solute form solid solutions and association of the solute does not occur. Such cases have been excluded from the tables, but aniline, pyrrole, pyridine (pp. 143, 144), and cyclohexanone and its derivatives (p. 146) in cyclohexane or dicyclohexyl are doubtful. Probably association has little, if anything, to do with the abnormal numbers found.

A considerable number of freezing-point and boiling-point measurements on aqueous solutions, mainly of inorganic substances, are given in Landolt-Börnstein, *Physikalisch-Chemische Tabellen*, to which source the reader is referred.

ALCOHOLS.

M.W. (Obs.).	43-144	32.0-31.6 73-166 80.4-214'9	79°8-178°3	93-144	102-143	116-158 118-169 121-151 131-185	122-272 108-134 144-166 111-161 132-178 130-194 238-265
Concentration.	0.32-7.29	3.35-10.92 0.220-10.43 0.1785-8.351	o'5325 11'943 o'217-11'19	0.243-10.07	0.248-10'14	0'257-12'16 0'248-11'65 0'263-10'95 0'325-12'44	0.273-17.18 0.346-8'586 1'84-11'73 0'267-12'34 0'257-12'48
Solvent.	Benzene ¹ , ² Nitrobenzene ²	Water 141 Benzene 3, 1, 6 Bromoform 4	Nitrobenzene v Benzene ³	Benzene 3	Benzene 3	Benzene 3 Benzene 3 Benzene 3 Benzene 3 Benzene 3	Benzene 3
M.W. (Calcd.).	32	74	88	100	102	114 128 130 142	130 116 130 128 144 156
Formula.	СН3ОН	(СН ₃₎₃ С.ОН	(CH ₃) ₂ C.OH	С. Н. (С. Н.). (С. Н.). (С. Н.). (С. Н.). С. Н.	Сон	(CH ₃) (C ₃ H ₃) (CH ₃) (C ₃ H ₃) (C ₃ H ₃) C.OH (CH ₃) (C ₃ H ₃) (C ₃ H ₃) C.OH (CH ₃) (C ₄ H ₃) C.OH (CH ₃) (C ₄ H ₃) C.OH (CH ₃) (C ₃ H ₃) C.C (CH ₃) ₃	$(CH_3) (C_9H_{13}) CHOH (C_2H_{5/2} (C_2H_{13}) CHOH (C_2H_{5/2} (C_3H_{1}) COH (C_2H_{5/2} (C_3H_{1}) COH (C_2H_{5/2} (C_3H_{1}) COH (C_2H_{5}) (C_3H_{1}) COH (C_3H_{5}) (C_3H_{1})_2 COH (C_3H_{5/2} (C_3H_{1})_3 COH (C_6H_{5/3} (C_3H_{1})_3 COH (C_6H_{5/3} (C_3H_{1})_3 COH (C_6H_{5/3} (C_6H_5) (C_6H_{5/3} (C_6H_5) (C_6H_5)$
Substance.	Methyl alcohol (carbinol)	Trimethyl carbinol	Dimethyl ethyl carbinol	Dimethyl allyl carbinol	Methyl diethyl carbinol	Methyl ethyl allyl carbinol Methyl propyl allyl carbinol Methyl dipropyl carbinol Methyl n-butyl allyl carbinol Methyl allyl tertiary-butyl car-	Methyl hexylcarbinol Triethyl carbinol Diethyl propyl carbinol Diethyl allyl carbinol Ethyl dipropyl carbinol Allyl dipropyl carbinol Ariphenyl carbinol

APPENDIX

Diphenyl carbinol (Benz-	$(C_6H_5)_2$ CHOH	184	Benzene 8	1.065-20.20	181-271
nydroi) Ethyl alcohol	C_2H_5OH	46	Ethylene bromide 7 Methyl succinate 7	0.294-3.633	49'2-134'3 46'1-58'7
			Veratrol' Benzene 6, 10, 8, 2	0.443-5.003	49.7-09.2 63.84-248.4
			Acetic acid ', * Water ¹⁰⁸	0.575-570	4/-39
			p -Bromotoluene 9 Bromoform 4	0.259-40.57	49.2-49.1 55.9-408.0
			Nitrobenzene 5, 2	0.4047-46.288	54-456
n-Propyl alcohol	C_3H_7OH	9	Dicyclohexyl 124	0.85-3.97	125.7-277.1
iso-Propyl alcohol	C ₃ H ₇ OH	9	Denzene 5, 5, 5 Nitrobenzene 5	0.2686-52.466	57-583
	110 110	i	Benzene 3, 2	0.22-12.72	73-212
n-Butyl alcohol iso-Butyl alcohol	C4H9OH	74 74	Benzene 3, 6, 1	0.225-10.97	75-188
			Cyclonexane		1 - 16
secyButyl alcohol	C_4H_9OH	74	Benzene 3	0.234-12.01	75-181
	110	00	Cyclohexane 125	0.33-9.41	132 1-309 5
Amyl alcohol	C, H.: OH	88	Cyclollexalle Benzene 3	0.215-12.64	78-226
430-mily1 alcollo1	22011-20		Cyclohexane 125	0.66-8-36	173.5-384.3
Amyl mercaptan	$C_{5}H_{11}SH$	104	Naphthalene 11	0.87-7.04	105-117
n-Heptyl alcohol	$CH_3 (CH_2)_5 CH_2OH$	116	Benzene	0.257-17.00	136-307
n-Octyl alcohol		130	Benzene 6	1.5-25.2	110.5-2561
Capity around	C.H.OH	242	Benzene 1, 6, 3	1.31-4.05	229-434
Cetyl arconol	16-33	+	Naphthalene 3	0.565-13.53	260-294
Allyl alcohol	CH2: CH.CH2OH	. 58	Benzene 12	0.48-11.21	68.3-202
. Linalool	$(CH_3)_2C:CH CH_3$	154	Benzene 13	0.35-14.31	141-208
	$\dot{C}H_{z}$ — $\dot{C}.OH.CH:CH_{\mathtt{z}}$				
			$ extstyle{ heta} extstyle{ h$	0.47-9.18	176-192

M.W. (Obs.).	85:57-130°0 102'8-136'7 95'57-103'0 106-139 118-271	112-342 112'2-262'2 173-182 170-189 211-230 130-323 150-232 155-199	139-156 179'0-340'6 151-322 176-192 140-161	M.W. (Obs.).	95'9-135'7 93'2-101'2 96'5-106'9 100-107 106'8-129'6 96'5 98'9-113 240-316
Concentration.	1'3-6'6 2'1-10'5 1'9-8'0 0'99-7'35 0'92-20'70	0.620-59'299 0.441-12'514 100-90'06 0'91-8'34 0'15-0'54 0'292-16'46 0'959-27'62 1'61-15 06	1'93-4'17 0'22-4'75 0'60-23'93 0'47-9'18 1'30-3'31	Concentration.	0.252-4.176 0.378-4.123 1.115-5.156 0.338-36156 1.001-18.22 1.178-7.598 4.641 0.39-5.34 1.75-3.12
Solvent.	Benzene ⁶ Benzene ⁶ Benzene ⁶ Naphthalene ¹ Benzene ¹	Nitrobenzene ⁵ Bromoform ⁴ Bromoform ⁴ Naphthalene ¹¹ Benzene ³	Ether ¹⁵ Cyclohexane ¹²⁵ Cyclohexane ¹²⁵ Benzene ¹³ , ¹⁶ p-Dichlorobenzene ¹³ Ether ¹⁵ Cyclohexane ¹²⁵	Solvent,	Ethylene bromide 7 Methyl succinate 7 Water 14, 10, 2 Benzene 8, 1, 10, 2 Acetic acid 8 Diphenylmethane 29 Ether 18 Naphthalene 19, 20 Cyclohexane 21
M.W. (Calcd.).	110°5 129 92°5 108	153 153 212 134 154	156	PHENOLS. M.W. (Calcd.).	46
Formula.	CH,CLCHOH: CH,OH CH,CLCHOH.CH,CL C,H,OCI C,H,CH,OH	C_6H_4 (NO ₃) CH_2OH C_6H_3 (NO ₃) CH_2OH C_6H_3 , $CHOH$, CO , C_6H_5 C_6H_5 , CH , CH , CH_2OH $C_{10}H_{18}O$	$C_{10H_{20}O}$	PHE Formula.	С ₆ Н ₅ ОН
Substan.	Monochlorhydrin Dichlorhydrin Epichlorhydrin Benzyl alcohol	o-Nitrobenzyl alcohol p-Nitrobenzyl alcohol Benzoin Cinnamic alcohol Borneol	Menthol	Substance	Phenol

238'9 117'6:349'6 98'7-252'6 85-127 123-136 138-151 144-153	145-157 132-145 147-194 172-253 171-219	151-194 156-196 270-253	135-158 161-170 161-170 158-182 158-182 173-213 156-227 168-252 172-353 133-210	136-138 300-310 186-177 170-185	193-188 219-257 212-252 228 210-238 220-231
1.32 2.179-42-877 0.1987-9.10 1072-17-25 0.93-5-39 0.93-8-89 0.93-8-89 0.93-13-86	0.72-5'99 0.84-5'66 0.82-12'08 1'04-5'34 0'93-3'88	3.052-7.134 2.203-5.641 0.88-8.48	0.93-6.86 0.92-7.66 1.04-7.28 0.86-6.15 0.88-6.15 0.50-2.45 0.88-6.59	0.411-1439 0.67-12'94 0.46-1'75	0.38-2.78 2.29-10.50 1.004-6.673 0.865 0.34-2'81
Dicyclohexyl 121 \$\rho\$-Bromotoluene \(^9\) Bromoform \(^4\) Nitrobenzene \(^5\) Nitrobenzene \(^2\) Nitrobenzene \(^2\) Naphthalene \(^2\) "Dinitrobenzene \(^2\)	p-Bromobenzene 22 Benzil 23 Naphthalene 19, 33 p-Dibromobenzene 22 p-Chlorobromobenzene 22 p-Chlorobromobenzene 22	p-Dichioobelizene Benzene 17 , 1 Water 17 Nitrohene 22	m-Dinitrobenzene 22 4-Dinitrobluene 22 2-4-Dinitrophenol 22 4-C Traitrophenol 22 5-Dichlorobenzene 22 6-Chlorobromobenzene 22 7-Chlorobromobenzene 23 7-Chlorobromobenzene 24 7-Chlorobromobenzene 24 7-Chlorobromobenzene 24 7-Chlorobromobenzene 24	p-Azoxyanisole 24 Naphthalene 19 Nitrogen peroxide 26 Northhalene 19, 23	Benganan Britagen peroxide 26, 134 Nitrogen peroxide 27 Chloroform 27 Acetic acid 28 Benzene 13, 27, 1 Naphthalene 19
139	139	139		297 184	184 229
$\mathrm{C_6H_4(NO_2)OH}$	$C_6H_4(NO_2)$ (OH)	$\mathrm{C_6H_4(OH)NO_2}$		$C_6H_2\ (Br)_3\ (NO_3)OH \ C_6H_3\ (OH)\ (NO_2)_2$	C ₆ H ₂ (OH) (NO ₂) ₂ C ₆ H ₂ (OH) (NO ₂) ₃
o-Nitrophenol	m-Nitrophenol	ho-Nitrophenol		oro-Dibromo \$\rho\$-nitrophenol 2'4-Dinitrophenol	a-Dinitropheno 2'4'6-Trinitrophenol

Substance.	Formula.	M.W. (Calcd.).	Solvent.	Concentration.	M.W. (Obs.).
o-Cyanophenol	C ₆ H ₄ (OH) CN	611	ϕ -Dibromobenzene ²² f -Chlorobromobenzene ²³ f -Dichlorobenzene ²³	1.03-511	152-310
			p-Bromotoluene 13	66.1-2.0	179-314
			p-lodotoluene 13	0.47-1.59	206-317
			p-roludine 13	0.86-6.78	144-109
•			Methyl p-toluate 13	1.84-7.77	120-136
			Methyloxalate 24	60.6-66.0	119-138
			Naphthalene 22	2.19-18-25	149-342
m-Cyanophenol	C ₆ H ₄ (OH) CN	611	p-Dibromobenzene 22	1.06-5.08	165-321
			p-Chlorobromobenzene	0.74-3.00	197-293
			p-Dichlorobenzene 22	1.56-7.63	194-421
			p-Xylene 13	0.40-1.30	143-190
			p-Bromotoluene 13	0.23-1.97	173-264
	•		p-Iodotoluene 13	0.48 - 1.70	194-308
			p-Nitrotoluene 13	1.060.1	142-163
			Methyl p -toluate 13	1.58-10.98	133-138
	C ₆ H ₄ (OH) CN	611	ϕ -Dibromobenzene 22	0.12-3.06	176-349
			p -Chlorobromobenzene 22	0.26-3.85	175-427
			p-Dichlorobenzene 22	0.74-2.28	195-305
			p-Nitrotoluene 13	1.14-6.25	174-187
			p-Toluidine 13	18.8-11.1	110-114
			Methyl p -toluate 13	2.27-10.92	123-138
			Naphthalene 25	00.01-66.0	131-248
			Methyloxalate 24	28-8-86.0	127-139
3-Nitro p-cyanopheno	C_6H_4 (OH) (CN) NO_2	164	Naphthalene 25	1:05-0.1	163-171
3.5-Dichloro p-cyanophenol	C_6H_2 (OH) (CN) CI_2	188	Naphthalene 25	21.9-26.0	188-208
3-Bromo p-cyanophenol	C ₆ H ₃ (OH) (CN) Br	198	Naphthalene 25	1.02-7.80	201-233
3.5-Dibromo p-cyanophenol	C ₆ H ₂ (OH) (CN) Br ₂	277	Naphthalene 25	1.34-8.23	277-293
3.5.Diiodo p-cyanophenol	C_6H_2 (OH) (CN) I_2	371	Naphthalene 25	1.28-6.14	366-370
o-Chlorophenol	C ₆ H ₄ (CI) OH	128.5	Naphthalene 19	1.29-8.74	123-137
p-Chlorophenol	C_6H_4 (CI) OH	128.5	Naphthalene 19, 20	0.45-10-21	137-157

174-186 161-187 7 153-196 313-8-327 313-8-328		A A A E.	122	35 186-208 17 275-285 17 275-285 18 26-372 11 202-316 14 184-203 14 284-354 16 228-240 17 239-307 18 39-307 18 39-307 19 39-307 19 30-478
1.54-9.66 0.58-9.46 0.47-10.27 2.5-3.99 0.46-2.09 1.48-6.84	0.37 0.37 1.09-8-24 1.23-10.11 1.08-9:57 1.08-9:71 1.08-9:11	1.20-10'21 1.21-9'85 1.12-6'90 1.15-8'75 0'95-7'19	1.15-10.30 0.36-10.14 0.411-38'96 0.63-9'62 0.5863-23'43 3.520-24'16 0.73-13'06	2'192-11'335 2'148-10'1092 1'840-8'887 0'792-5'011 0'888-4'734 1'584-4'724 1'584-4'724 1'253-4'697 2'133-10'122
Naphthalene ^{19, 20} Benzene ¹³ Naphthalene ¹⁹ Benzene ¹ Cyclohexane ¹²⁸ Naphthalene ¹⁹	Naphthalene ²⁰ Benzene ¹ Naphthalene ²⁰	Naphthalene ²⁰ Naphthalene ²⁰ Naphthalene ²⁰ Naphthalene ²⁰ Cyclohexane ¹²⁵ Dyclohexane ¹²⁵	Denzene 19, 33 Paphthalene 19, 33 \$\$P\$-Bromotoluene 9 Cyclohexane 135 Nitrobenzene 5 Stearic acid 29 Cyclohexane 125 Cyclohexane 125	Benzene 30 Benzene 30 Benzene 30 Benzene 30 Panzene 30 p -Dibromobenzene 30
173	243 243 243 243 243 228	242 242 108	108	187 266 266 266 345 187'5 187'5 180 232 232 232 232 232 232 232 232 232 23
C ₆ H ₄ (OH) Br C ₆ H ₄ (OH) Br C ₆ H ₂ (Br) ₃ OH	C ₆ H ₃ (CH ₃) C ₆ H ₃ (N; NC ₆ H ₃ OH C ₆ H ₄ (NO ₂).N: N.C ₆ H ₄ OH C ₆ H ₄ (NO ₂).N: N.C ₆ H ₄ OH C ₆ H ₄ (NO ₂).N: N.C ₆ H ₄ OH C ₆ H ₄ (NO ₆).N: N.C ₆ H ₄ OH C ₆ H ₄ (NCH ₃).N: N.C ₆ H ₄ OH C ₆ H ₄ (NCH ₃).N: N.C ₆ H ₄ OH C ₆ H ₄ (NCH ₃).N: N.C ₆ H ₄ OH	C,H, (OC,H,),N:N.C,H,OH C,H, (OC,H,),N:N.C,H,OH C,H, (OC,H,),N:N.C,H,OH C,H, (OC,H,),N:N.C,H,OH	C_6H_4 (CH ₃) OH .	C ₆ H ₃ (OH) (CH ₃) Br C ₆ H ₃ (OH) (CH ₃ Br) Br C ₆ H ₃ (OH) (CH ₃ Br) Br C ₆ H ₃ (OH) (CH ₃ Br) Br C ₆ H ₃ (OH) (NO ₂) CH ₂ Cl C ₆ H ₃ (OH) (NO ₂) CH ₂ Cl C ₆ H ₃ (OH) (NO ₃) CH ₂ Cl C ₆ H ₃ (OH) (NO ₃) CH ₂ Br C ₆ H ₃ (OH) (CH ₃) CO ₂ C ₂ H ₃ C ₆ H ₃ (OH) (CH ₃) CO ₂ C ₂ H ₃ C ₆ H ₃ (OH) (CH ₃) CO ₂ C ₂ H ₃ C ₆ H ₃ (OH) (CH ₃ Br) (CO ₂ C ₂ H ₃
o-Bromophenol f-Bromophenol Tribromophenol	f-Iodophenol Benzeneazophenol o-Tolueneazophenol m-Nitrobenzeneazophenol f-Nitrobenzeneazophenol o-Methoxybenzeneazophenol	p-thernoxypotrantazophono o-Ethoxybenzeneazophenol m-Ethoxybenzeneazophenol p-Ethoxybenzeneazophenol o-Cresol	<i>∮</i> -Cresol	p-Bromocresol p-Bromo-o-bromocresol 2-q-Dibromo-o-cresol 2-q-Dibromo-o-Bromocresol p-Nitro-p-chlorocresol p-Nitro-p-bromocresol p-Nitro-p-bromocresol p-Nitro-p-bromocresol p-Carbethoxy-o-cresol p-Carbethoxy-o-cresol p-Carbethoxy-o-chlorocresol p-Carbethoxy-o-chlorocresol p-Carbethoxy-o-chlorocresol p-Carbethoxy-o-chlorocresol

Concentration. M.W. (Obs.).	249-267 318-352 200-273 157-159 159-173 201-205 131-137 170-177 126-157 117-145	149-174 127-147 158-184 158-144 141-135	255-292 193-218 167-194	177.3-201.6	174.7-176.4 176.3-177.7	174'0-175'6 176'2-175'6
Concentration.	1.840-6.870 1.923-6.165 0.693-4.972 0.44-8.20 0.937-2.768 1.082-6.224 1.44-7.66 1.27-9.18 0.85-9.98 0.65-10.12 0.95-13 0.4	1.51-6.38 0.26-4.99 0.60-5.84 0.93-9.68 2.118-3.584 1.734-7.886	0.32-3.62	2.46-3.74 2.56-4.48	3.20-5.93 2.5-3.8	1'94-3'20 1'94-3'41
Solvent.	p-Dibromobenzene 30 p-Dibromobenzene 30 p-Dibromobenzene 30 Naphthalene 19 p-Dibromobenzene 30 p-Dibromobenzene 30 p-Dibromobenzene 30 Naphthalene 11 Naphthalene 19 Naphthalene 19 Naphthalene 19 Naphthalene 19 Naphthalene 19	p-Dibromobenzene ³¹ Benzene ¹ p-Dibromobenzene ³¹ Pyridine ³² Benzene ³³ , ¹ Stearic acid ²⁹	p-Dibromobenzene ³¹ p-Dibromobenzene ³¹ p-Dibromobenzene ³¹	Acetone 34 Ethyl alcohol 34	Acetone 34 Ethyl alcohol 34	Acetone ³⁴ Ethyl alcohol ³⁴
M.W. (Calcd.).	232 311 153 153 153 153 154 124 122 122	144	189 189 189	173		
Formula.	C,H. (NO.) (Br) (OH) (CH.) C,H. (NO.) (Br) (OH) CH.,Br C,H. (NO.) (Br) (OH) CH.,Br C,H. (CH.) (NO.) OH C,H. (CH.) (CH.) SH C,H. (CH.) (CH.) OH C,H. (CH.) (CH.) OH C,H. (CH.) OH	$C_{10}H_8^{10}$	$C_{10}H_6$ (NO ₂) OH	ON HOW		HON
Substance.	o-Bromo-p-nitro-o-cresol o-Bromo-p-nitro-o-cresol p-Nitro-acetoxycresol m-Nitro-p-cresol p-Nitro-p-cresol p-Nitro-p-cresol p-Thiore-cresol p-Thiocresol o-Methoxy-m-thiocresol o-Xylenol p-Xylenol p-Xylenol p-Xylenol p-Xylenol	a-Naphthol B-Naphthol	p-Nitro-a-naphthol B-Nitro-a-naphthol a-Nitro-B-naphthol	Nitroso-naphthol		

Decahydronaphthol Pseudo-cumenol Guaiacol Thymol	$C_{0,H_{18}}^{1,9}O \\ C_{6,H_{2}}^{1}(CH_{3}^{1,8})OH \\ (CH_{3}^{1,8},C_{6}^{1}H_{2}^{1}OH \\ (CH_{3}^{1,2},CH,C_{6}^{1}H_{3}^{1}(CH_{3}^{1})OH \\ \end{array}$	154 136 124 150	Naphthol ¹²⁸ Naphthalene ²⁰ Naphthalene ¹⁹ , ²⁴ Ethyl ether ¹⁴ , ^{1b} Benzene ¹ Naphthalene ¹⁹	0.47-2'94 1'58-9'85 0'70-9'07 2'35 1'28-17'71	148-141 138-156 114-135 145 146-184
Eugenol	C ₆ H ₃ (C ₃ H ₅) (OCH ₃) OH	164	\$\frac{\}{\}\cdot \cdot	0.369-13'057 0'1913-10'16 0'4628-30'18 0'37-14'94 0'995-5'997	132'8-171'6 121'7-193'9 124-187 140'1-395'8 175 193'5
	DIHYDROX	DIHYDROXYPHENOLS.			
Substance.	Formula.	M.W. (Calcd.).	Solvent,	Concentration.	M.W. (Obs.).
Catechol	$\mathrm{C}_6\mathrm{H}_4(\mathrm{OH})_2$	011	Water 17 Benzene 17, 1	3.283-5.218 3.085-6.639	123.9-127.4
			Etner 13 Naphthalene 19, 24	2.34 0.46-6.42	116-144
Resorcinol	$C_6H_4(OH)_2$	011	Water 17 Benzene 17 Benzine 32	3.286-3.997	119.7-119.9
			Ether 37, 36, 33 Ethyl alcohol 36, 37, 38 Naphthalene ¹⁹ , ²⁴	2.06-8.99 1.99-5.05 0.44-2.88	124-105 112-127 110-5-113'0 112-139
Quinol	$C_6H_4(OH)_3$	011	Acetone 38 Ether 18 Naphthalene 19	5.81 1.913 0.36	98 113 90
Resorcinol methyl ether	ОН	124	Naphthalene 24	0.55-6.78	125-152
Hydroquinone methyl ether	OH OCH3	124	Naphthalene 24	60.E-80.I	127-144
Styphnic acid	$C_6H(NO_2)_3$ (OH) ₂	245	Naphthalene 19	969.0	259

TRIHYDRIC PHENOLS.

M.W. (Obs.).	128-143 130	139-142 168-156		M.W. (Obs.).	1.66	407-511 194-178 119-128 121-7-140:0	133.2-166.6 76.2-94.1	60.6-77.4 63.8-84.1	82.0-250.0 81.3-174.6	85-157	94.9-124.4	9477-1123 246-278 80-99 183-225 209-231
Concentration.	0.31-1.04 1.85	5.540-12.79 4.635-10.98		Concentration.	0.849-34.96	0.28-2/25 0.28-2:25 0.46-1:44 0.86-3.40	0.75-3.03	0.421-3.334	0.345-14.00 0.3395-34.74 0.2240-7.40	0.346-18:10	2.83-14.82	0.90-7'00 0'98-9'64 1'05-4'77 5'038-9'124 3'556-8'740
Solvent.	Naphthalene 19 Ethyl ether 14	Water ¹⁷ Water ¹⁷		Solvent.	Nitrobenzene 5 Benzene 2 W. A. A. S.	Vyater Nitrogen peroxide ¹³⁴ , ²⁶ Carbonyl chloride ³⁹ Cyclohexane ¹²⁵	Dicyclohexyl 124 Sulphur dioxide 39 Fthylane bromide 7 2	Methyl succinate? Veratrol 10 8 9	Denzene v, v, *. p-Bromotoluene 9 Bromoform 4	Nitrobenzene ⁵ , ² Water ¹⁴⁹	Phosphorus oxychloride 126 Benzene 1	Frosphorous oxycnioride for Benzene 1 Naphthalene 11 Water 17 Benzene 17, 1
M.W. (Calcd.).	126	126		M.W. (Calcd.).	46	9					94.5	139 76 136
Formula.	C_6H_3 (OH) $_3$	S ₆ H ₃ (OH) ₃	ACIDS.	Formula.	н.соон	сн.,соон					CH ₂ Cl.COOH	CH,Br.COOH CH,COSH C,H,CH,.COOH
Ľ.	ເ ຶ່ວ	ິ່ວ			д	Ċ					СН	CH ₃ CH ₄ C
Substance.	Pyrogallol	Phloroglucinol		Sub tance,	Formic acid	Acetic acid					Chloroacetic acid	Bromoacetic acid Thioacetic acid Phenylacetic acid

Phenoxyacetic acid Dichloroacetic acid Trichloroacetic acid	С ₆ н ₅ о) сн. соон СНСІ ₃ -соон ССІ ₃ -соон	152 128'9 163'5	Naphthalene ²⁰ Ethylene bromide ¹⁴ Phosphorus oxychloride ¹²⁶ Formic Acid ⁴¹ Phosphorus oxychloride ¹⁷	0.62-9.15 0.57-4.93 0.69-8.07 1.367-2.876 0.94-10.88	204-254 168-210 115'9-117'3 158'6-139'0 142'9-134'6
Tribromoacetic acid	СВг. СООН	297	Nitrogen peroxide ¹⁹³⁴ Benzene ¹ Nitrogen peroxide ²⁶	1.00-1.77 1.00-5.56 0.93-3.80	258 ³ -277 ³ 312-467 552-606
Propionic acid	C_2H_b .COOH	74	Formamide 43 Water 17 Denoted 17	3.359-4.748 0.823-15.47	242'9-237'4 66'8-84'8
Phenylpropionic acid	C ₆ H ₅ ·CH ₂ ·CH ₂ ·COOH	150	Phosphorus oxychloride 126 Dinhenyl 29	0.81-7.09	97.1-128.1 97.1-128.1 260.8
			p-Cresol 29 Anethol 29 Azoberzene 29	2.605-11.40 1.710-17.46	159'2-164'5 258'1-236'2
Butyric acid	С3Н,.СООН	88	Phosphorus oxychloride 126 Water 17 Nitrogen peroxide 134	0.64-4.94 1.847-9.605 0.507-1.22	81.7-99.1 81.7-99.1 208-217
				0.951-11.20 0.6924-11.2364	157.0-174.6
Trichlorobutyric acid	C ₃ H ₄ Cl ₃ .COOH	2.161	Nitrobenzene 5 Nitrogen peroxide 26 Bargan 1	0.3330-37.3474 0.85-6.97	104-259 233-281 333
Valeric acid	СН3(СН2)3СООН	102	Nitrobenzene 5, 2	0.3750-41.49	353 143-351
Isonitrosovaleric acid	CH ₃ .CH(NO)(CH ₂) ₃ COOH	131	Denzene - Phenol 43 Benzene 1	1.575-9799	139-203.8
ਾਰ	CTHIS COOH	144	Benzene 1	65.11-66.1	247-255
	CH ₂ Cl.CH: CH.COOH	200	Denzene 1 Benzene 1	0.72-10.30	320-370 164-227
Carvocrotonic acid	С,Н,СН:СН.СООН	190 148	Benzene 1	1.04 1.45	308 258
Phenylcinnamic acid	C ₆ H ₅ CH: C.(C ₆ H ₅).COOH	224	Nitrobenzene ^{44, 5} Diphenylmethane ²⁹ Phenol ²⁹ Benzene ¹	0.44-1'20 0'389-1'559 7'355-9'638 0'95	265-307 331'6-382'9 194'8-196'1 348

Substance,	Formula.	M.W. (Calcd.).	Solvent,	Concentration.	M.W. (Obs.).	
Oxalic acid Malonic acid	COOH.COOH CH ₂ (COOH),	90 I04	Water 17 Ethyl Ether 14	4.700-9.677	72.1-74.7	
Succinic acid	$(CH_2,COOH)_2$	811	Ethyl alcohol 14	2.12-5.78	123-113	
			Water 17	4.487-7.591	123-118'9	
			Pyridine 32	1.18-8.40	127-113	
Denzoic acid	C ₆ H ₅ .COOH	122	Nitrogen peroxide 28	2.42-9.56	209-228	
			Carbon disulphide 14, 46	60.6-2.1	237-273	
			p-Dibromobenzene 13	0.47-4.14	196-255	
			m-Dinitrobenzene 13	1.12-6.17	148-190	
			Methyl p -toluate 13	0.20-8.68	163-170	
			Carbonyl chloride 39	19.1-65.0	244-267	
			Sulphur dioxide 39	1.47-7.85	214-237	
			Nitrobenzene 44, 2	0.305-2.01	260-273	
			Water 17	6.088-7.288	235-258	
			Benzene 14, 8, 38, 2	2.61-10.75	210.250	
			Acetic acid 8, 14	0.857-24.92	123-146	
			Ether 14, 18	1.26-22.10	120-148	
			Phenol 47, 29	0.27-5.08	126-138	
			Chloroform 14, 38, 48	0.85-8-61	199-226	
			Ethyl alcohol 14, 38, 48	1.98-4.42	115-124	
			Ethyl acetate 14	0.79-18.55	121-144	
			Naphthalene 20	59.6-61.I	194-232	
			Phosphorus oxychloride 126	0.73-1.79	139.0-154.8	
			Acetone 38	8.58-4.97	115	
			p-Toluidine 29	2.314 9.139	158.3-139.8	
1		•	Cyclohexane 125	0.45-0.55	227.6-231.1	
0-Cillorobenzoic acid	Cent.COOH	156.5	Benzene 1	0.28	258	
1			Naphthalene 20	0.22-5.62	199-261	
m-Cilloropenzoic acid	C.H.C.COOH		Naphthalene 20	0.28-6.94	212-280	
o-bioliopelizoic acid	CeH4BI.COOH	201	Naphthalene 20	0.57-9.13	269-340	
M. Bromobengoio coid	11000 -0 11 0		Benzene *3	2.47-19.4	307-364	
m-Dionicoenzoic acid	Centbr.COOH	201	Bromine 49	0.29-0.81	290-30I	
A. Bromohennois soid	11000 4110		Naphthalene 20	0.26-8.86	287-367	
ל-טוווסטפווקסוכ מכות	C ₆ H ₄ Br.COOH		Naphthalene 20	0.43-0.73	283-313	

0.59-9'00 320-417 0'51 210 0'64-5'77 201-249	0.56 211 0.28-8.86 214-310 0.45-1.40 145'8-121'0	0.34-8.69 239-306		0.39-150 277-301 0.58-5:80 175-230 0.44-8:10 200-242		0.62-6.54 209-252 1:01-1:86 194-197	6		1.399-24 43 1.59 1.69 2.497-4.344 192'6-187'4		Concentration, M.W. (Obs.).	1.81-12.10 160-165 1.67-21.46 157-147	٠
Naphthalene 20 Senzene 1 Naphthalene 20	Naphthalene 20 Naphthalene 20 Phosphorus, oxychlor 1e 126	Naphthalene 20 Renzene 1	Naphthalene ²⁰ Naphthalene ²⁰ Naphthalene ²⁰	Naphthalene 20 Naphthalene 20 Nitrogen peroxide 26	Naphthalene 20 Nitrogen peroxide 26	Naphthalene 20 Nitrogen peroxide 26	Water 17	Acetic acid 50 Benzene 8, 14	Acetic acid ϕ . Acetic acid ϕ -Toluidine		Solvent.	Pyridine 32 Fthyl alcohol 14, 33, 38	Pyridine ³² Water ¹⁷ Chloroform ¹⁴ , ⁸⁸
248 137	137 167		212	257 136	136	136	991	111	148	HYDROXY ACIDS	M.W. (Calcd.).	150	138
C ₆ H ₄ I. COOH C ₆ H ₄ (NH ₂). COOH	$C_6H_4(NH_2).COOH C_6H_4(NO_2).COOH$	C ₈ H ₄ (NO ₂).COOH	C ₆ H ₄ (NO ₂).COOH C ₆ H ₃ (NO ₂) ₂ .COOH C ₆ H ₄ (NO ₃) ₂ .COOH	C,H2 (NO2)3COOH C,H4 (CH3).COOH	С6Н4 (СН3).СООН	C ₆ H ₄ (CH ₃).COOH	C ₆ H ₄ (COOH) ₂	C,H,NH,COOH (C,H,CO),0	C ₆ H ₄ (CO) ₂ O	HYDROX	Formula.	соон.снон.снон.соон	С,н, (он),соон
o-Iodobenzoic acid Anthranilic acid	\$-Amidobenzoic acid o-Nitrobenzoic acid	m-Nitrobenzoic acid	p-Nitrobenzoic acid o-o-Dinitrobenzoic acid Pinitrobenzoic acid	sym-Trinitrobenzoic acid	m-Toluic acid	p-Toluic acid	Phthalic acid	a-Carboxypyrrolic acid Benzoic anhydride	Phthalic anhydride		Substance.	Tartaric acid	Salicylic acid

M.W. (Obs.).	165-234 129-143 137-136 179-258	207.0-159.2	218 141-157	179'1 185'8 155-159	209-260 188-193		M.W. (Obs.).	153-197			139-201.0 167-170		274-310		384-394	
Concentration.	0.75-5'61 1.37-17'29 1.40-2'86 0.52-8'19	4.864-7.986	0.21 4.318-7.236	1.252-3'430 5'137-6'908	2.263-4.063 5.556-11.46		Concentration.	1.11-5.49	0.22-7.93	0.946-3.403	0.76-5.58		0.43-2.86		0.46-1.17	
Solvent.	Benzene 14, 1, 38 Ethyl alcohol 14, 53, 38, 54, 48 Ether 14, 53 Naphthalene 20 4. Tolvising 29	Water 17	Naphthalene 20 Water 17	<i>p</i> -Toluidine ²⁹ Water ¹⁷	Benzene ¹⁷ Water ¹⁷		Solvent	Benzene 12 Benzene 1	Benzene ⁱ Benzene ¹	Ethyl alcohol ⁵⁹ Benzene ¹	Cyclohexane 127 Alcohol 59		Benzene 12		Benzene 12	
M.W. (Calcd.).		138	138	152	154	RS.	M.W. (Calcd.).	160 188	130 129	68 130	152		286		348	
Formula.		C_6H_4 (OH).COOH	С ₆ Н ₄ (ОН).СООН	С ₆ Н ₅ СН (ОН).СООН	C_6H_3 (OH) $_2$ COOH	ESTERS	Formula.	COOCH, CH, CO.COOCH, COOC, H, CH, CO.COOC, H,	CH3.C (NH ₂): CH.COOC ₂ H ₅	NaOC2H5 CH3,CO.CH2,COOC,H5	CH3.C(ONa) : CH.COOC,H5	cH.	CH, CO.CH.CH.CH.CO.CH,	COÓC2H, COOC2HS	сн.со.сн.сн.со.сн.	COOC,H, COOC,H,
Substance.		m-Hydroxybenzoic acid	p-Hydroxybenzoic acid	Mandelic acid	Protocatechuic acid		Substance.	Dimethyl oxal-acetate Diethyl oxal-acetate Ethyl formyl monitarie	Ethyl 6-amido-crotonate	Sodium etnylate Ethyl acetoacetate	Monosodium ethyl acetoacetate		Ethylidene diacetoacetate		Benzylidene diacetoacetate	

98'9-162 107-176 160-177 155-204 172'4-243'2 180'5-229'9 207'4-221'7 190'8-172'2		. 44 444	2014-215-7 185-348 232-5-219:3 266-244-8 209-6-183:2 2127-194:5 220-6-305:1	228-354 194-8-204-2 171-8-196-2 205-1-226-6 182-8-229-5 2100-344-2
0.70-9:28 0.40-11:82 1:234-5:782 2:082-6:127 1:50-8:08 1:32-13:26 1:97-18:48 1:55-16:32	0.80-10'49 3.03-12'62 2.06-9'23 4'31-15'71 4'51-18'20 crys, 2.10-12'70	3.13-14'09 3.13-14'09 4.14-23'10 3.84-22'07 1.08-17'55 1'26-16'99	1.91-900 1.23-23.10 1.38-6-29 2.12-9.14 3.57-12-85 5.48-12-64 3.64-12-85	2.78-8.40 1.80-14.36 4.54-10.58 4.54-10.98 4.61-10.90 0.386-7.895
Benzene 12 Benzene 12 Ethyl alcohol 59 Ethyl alcohol 59 Benzene 60 (cryos.) Benzene 60 (chul.) Methyl alcohol 61, 60 Ethyl acetate 60 Acetone 60, 61	Chloroform ⁶⁰ Ethyl acetate ⁶⁰ Chloroform ⁶⁰ Methyl alcohol ⁶⁰ Acetone ⁶⁰ Benzene ⁶⁰	Acetone 60 Methyl alcohol 60 Ethyl acetate 60 Chloroform 60 Carbon disulphide 60 Benzene 60 (ebul.)	Benzene ¹⁰ (cryos.) Benzene ¹⁵ Ethyl accetate ⁶⁰ Chloroform ⁶⁰ Methyl alcohol ⁶⁰ Accetone ⁶⁰ Benzene ⁶⁰ Benzene ⁶⁰	Ethylene bromide ⁶² , ⁶³ Water ⁶⁴ Methyl alcohol ⁶³ , ⁶⁰ , ⁶⁴ Ethyl alcohol ⁶⁴ <i>n</i> -Propyl alcohol ⁶⁴ Benzene ⁶⁵ (cryos.) Benzene ⁶⁵ (cryos.)
104 118 160 182 162'I	1.061	204	178	506
CH,CH (0H).COOCH, CH,CH (0H).COOC,H, CH,: (COOC,H,), CHNa: (COOC,H,), CHNa: (COOC,H,),	Соос <u>,</u> н ₅ .снон.сн ₂ .соос ₂ н ₅	сн (сосн ₃) сн (он) (соосн ₃₎₂	(СНОН.СООСН ₃)₂	$(\mathtt{CHOH}.\mathtt{COOC}_{\mathtt{2}}\mathtt{H}_{\mathtt{5}})_{\mathtt{2}}$
Methyl lactate Ethyl lactate Ethyl malonate Monosodium ethyl malonate Dimethylmalate	Diethylmalate	Dimethyl acetyl malate	Dimethyl tartrate	Diethyl tartrate

.51										-								
M.W. (Obs.).	205'5-539'4 259'5-209'9 242'4-217'1 235'4-234'8 274'3-318'5	252'5-307'4 258	208'7-226'1 135'7-193'6	159-163 128-151	156-157 159-147 151-171	142-149	145-156	168-302	224-362	161-236	194-260 193-200	219-312	152-353	771-771	178-318	151-148	149-167	175-193
Concentration.	0.296-16'75 3'81-17'32 0'91-11'50 2'00-9'91 5'07-16'92	2.51-11.10	3.52-22'41 1.01-19'19	1.54-6'27 0'56-1'92	0.95-1.97 0.53-1.36 1.66-23.12	0.625-1.755	0.02-5.34	0.97-4.95	1.88-8.77	00.8-00.1	0.78-3.53	0.35-2.12	0.18-3.88	2.55-10.05	1.26-14.72	0.50-1.415	1.01-9.02	0.18-1.80
Solvent.	p-Xylene 66 Acetone 60, 61 Ethyl acetate 60 Chloroform 60 Benzene 60 (ebul.)	Sulphur dioxide 66	Benzene ⁶⁷ Acetic acid ⁶⁷	2.4-Dinitrotoluene 22.4-6-Trinitrotoluene 22	ρ -Nitrotoluene ²² ρ -Chloronitrobenzene ²² Naphthalene ²² , ¹⁹ , ²⁰ , ²³	<i>p</i> -Azoxyanisol ²⁴ Nitrobenzene ²²	p-Dibromobenzene 22 Benzil 22	p-Chlorobromobenzene 22	P-Diction openizene Benzene 13	Naphthalene 23, 19	p -Aylene ϕ -Chlorotoluene 13	p-Bromotoluene 18	p-lodotoluene 13	Methyl 6-tolnate 13	Naphthalene 22, 23, 24, 25	p-Azoxyanisol 24	Dimethyloxalate 24	Nitrobenzene 221
M.W. (Calcd.).		2.292	218	152				152							153			
Formula.		(CHOH.COOC, H ₉), CH ₂ .(COOCH ₃)	C00C2H5.CH.(C00CH3)	С ₆ Н ₄ (ОН),СООСН ₃				С ₆ Н ₄ (ОН).СООСН ₃							С ₆ Н ₄ (ОН).СООСН ₃			4
Substance.	·	Di-isobutyl tartrate	Ethyl diacetyl glycerate (active)	Methyl salicylate				Methyl m -hydroxybenzoate							Methyl p-hydroxybenzoate			

146-178 164-210 156-200 153-191 159-202 205-264 167-217 152-178 194-257 186-224	231-271	299-314	203-249	294-323	402-406	191-204	211-221 219-243	179-340	163-211 171-175 163-175 164-174 152-240 138-4-139	239-264'9 133'5 141'6 144'7-144'8
1'02-6'18 0'90-7'87 0'90-6'47 0'90-6'12 0'0-170 0'28-0'78 1'28-7'24 1'28-7'24 1'28-7'24 1'28-7'80	0.81-10.24	1.09-6.54	1.23-10.30	1.20-11.18	1.23-9.76	0.97-9.14	0.82-7.38	0.65-7.00	0.45-7.22 0.57-9.71 0.76-11.24 0.62-9.93 0.54-8.76 2.41.3.96	0.131 0.189 0.220-0.305 0.280-0.351
m-Dinitrobenzene 22 p-Nitrotoluene 23 24-Dinitrotoluene 22 2-4.6-Trinitrotoluene 22 p-Chloronitrobenzene 22 p-Chlorobenzene 22 p-Chlorobenzene 22 p-Chlorobenzene 22 p-Diptomobenzene 22 Benzil 22 Naphthalene 19 Benzene 18	Naphthalene 24, 25	Naphthalene 24	Naphthalene 25	Naphthalene 20	Naphthalene 20	Naphthalene 25	Naphthalene ¹⁹ Naphthalene ¹⁹ , ²³	Naphthalene 19	Naphthalene 19, 28 Naphthalene 19 Naphthalene 19 Naphthalene 19 Naphthalene 19 Accin acid 135, 92 Record 135, 92	Methyl iodide ¹¹³ Methyl iodide ¹¹³ Ethyl iodide ¹¹³
166 214 186'5	231	278	221	310	404	161	211 211	191	166 166 166 166 166	234 109 137
C ₆ H ₄ (OH).COOC ₂ H ₅ C ₆ H ₄ (OH).COOC ₆ H ₅ C ₆ H ₃ (OH).CI.COOC ₆ H ₅	C6H3 (OH).Br.COOCH3	С6Н3 (ОН).І.СООСН3	С6Н2 (ОН).С12.СООСН3	C ₆ H ₂ (OH).Br ₂ .COOCH ₃	C ₆ H ₂ (OH).I ₂ .COOCH ₃	C ₆ H ₃ (OH).NO ₂ .COOCH ₃	C ₆ H ₃ (OH).NO ₂ .COOC ₂ H ₅ C ₆ H ₃ (OH).NO ₂ .COOC ₂ H ₅	C ₆ H ₃ (OH).NH ₂ .COOCH ₃	C ₆ H ₃ (OH).(CH ₃).COOCH ₃ C ₆ H ₃ (OH).(CH ₃).COOCH ₃	(CH ₃) ₄ P ₃ O ₇ (CH ₃) ₂ PO ₃ (C ₂ H ₃) ₃ PO ₃
Ethyl p-hydroxybenzoate Phenyl salicylate Methyl 3-chloro-p-hydroxyben-	Methyl 3-bromo-\$-hydroxyben-	Methyl 3-iodo-p-hydroxyben-	Methyl 3.5-dichloro-p-hydroxy-	Methyl 3.5-dibromo-p-hydroxy-	Methyl 3.5-di-iodo-p-hydroxy-	Methyl 3-nitro-p-hydroxyben-	zoare Ethyl m-nitro-salicylate Ethyl m-nitro-p-hydroxyben-	Methyl amido- p -hydroxyben-	Methyl 6-hydroxy o -toluate Methyl 9-hydroxy p -toluate Methyl 4-hydroxy m -toluate Methyl 2-hydroxy m -toluate Methyl 2-hydroxy m -toluate Methyl 4-hydroxy o -toluate Dimethyl sulphate	Methyl pyrophosphate Methyl hypophosphate Ethyl hypophosphate

OXIMES.

Substance.	Formula.	M.W. (Calcd.).	Solvent.	Concentration,	M.W. (Obs.).
Aldoxime	CH ₃ CH: NOH	59	Diphenylamine ²⁹ Benzophenone ²⁹ Stearin ²⁹	0.405-9'89I 0.33I-3'185 0'80I-8'264	84.48-155.1 71.83-84.05 56.06-50.16
			Azobenzene 29 $ ho$ -Toluidine 29 Cetyl alcohol	0.339-3.340 0.507-54.93	64.83-84.80 68.39-68.25 67.10
Acetoxime	$(CH_3)_2$ C: NOH	73	Acetic acid 8, 55 Benzene 8	0.284-20.23	71.5-77.3
Benzaldoxime	C ₆ H ₅ .CH: NOH	121	Water 8 Benzene 8 Acetic acid 8	0.960-1935 0.439-3970	76.5-88.3
o-Nitrobenzaldoxime \$P.Nitrobenzaldoxime Benzaldoxime ethylether	$C_6H_4(NO_2)$ CH: NOH $C_6H_4(NO_2)$ CH: NOH C.H. CH: NOH	166 166	Naphthalene ¹¹ Naphthalene ¹¹ Naphthalene ¹¹ Benzene ⁸	1.01-9.49 1.06-5.96 1.14-9.41	124-181 171-209 170-209
Acetophenoneoxime	C_6H_5 .C. (CH_3) : NOH	135	Acetic acid 8 Benzene 8, 14	0.833-28:10 0.625-40:59	130-162 132-166 169-279
			Acetic acid 8 Ether 18 Ethyl alcohol 14	1.517-23'81 2.21 1.55-12'97	134-158 137 129-154
Benzophenoneoxime	$(C_6H_5)_2$, $C: NOH$	197	Cyclohexane 120 Acetic acid 8 Bosson 8	0.49-2.34	219.5-258'3
o-Amidobenzophenoneoxime a-Benzilmonoxime b-Benzilmonoxime Carvoxime	NH, C, H, C (C, H,): NOH C, H, C: (NOH) CO.C, H, C, H, C: (NOH) CO.C, H, + \(\frac{1}{2} \) C, H, \(\frac{1}{2} \)	232 225 264 165	Denzene 1 Benzene 1 Benzene 1 Benzene 1 Cyclohexane 125	0.238-15.88 0.22-0.78 0.07-0.80 0.65-3.89 0.38-4.80	211-250 187-253 179-270 278-356 290'5-335'8
d-Camphoroxime	C10F118: NOF	107 167 167	Benzene 19 Cyclohexane 195 p-Dibromobenzene 13 m-Dinitrobenzene 13 Methyl p-toluate 13 Acetic acid 8 Benzene 1, 8 Benzene 1, 8	1'43-13'41 0'24-3'96 0'98-8'48 0'50-7'05 1'29-8'95 0'880-45'31	228-305 215'8-324'4 200-276 151-227 306-291 164-200 182-286 229-308

AMIDES.

			•																	7-
	M.W. (Obs.).) 62·6-72·9 80-98 174-178 115·7·107·7		156-207 132-134 143-151	154-167	148-218	189-285	188-279	79.6-80.7	76.3-84.4	95.7-110 4	112-155	97.4-104.6	9.5-36.8	9.5111.011	137-168	84-88-6	89.6-103.9	9.511-8.901	128-165 123-136
	Concentration.	31'2-74'3 (N) 62'6-72'9 0'92-2'84 80-98 6'250-7'597 174'178 22'6-50'7 (N) 115'7-107'	34'3-66'5 (N) 23 2-31'8 (N) 9'6-11'6 (N) 10'0-44'2 (N)	5.346-12.04 25.4-5.83 (N) 28.8-51.0 (N)	20.3-31.0 (N)	1.53-8.41	1.95-13.24	2.01-9.42 5.496-11.68	42.4-80.7 (N)	33.8-68.7 (N)	25.5-03.2 (N) 11.3-12.0 (N)	21.4-56.7 (N)	4.337-7.099	28.2-58.9 (N)	39.2-59.6 (N)	25.8-50.7 (N)	4.455-7.450 27:0-66:0 (N)	29.1-70-3 (N)	32.5-52.6 (N)	25.2-498(N) 5.034-9.198
	Solvent.	Acetone 56 Naphthalene 11 Water 57 Ethyl alcohol 56	Acetone 56 Chloroform 56 Ether 56 Benzene 56	Water ⁵⁷ Ethyl alcohol ⁵⁶ Acetone ⁵⁶	Chloroform 56 Benzene 12	Naphthalene 58	Naphthalene 58	Naphthalene % Water 57	Ethyl alcohol 56	Acetone 56	Ether 56	Benzene 56	Water 57 Fthyl gloshol 56	Acetone 56	Chloroform 56	Benzene 36	water % Ethyl alcohol 56	Acetone 56	Chloroform 56	Benzene ⁵⁶ Water ⁵⁷
ES.	M.W. (Calcd.).	59 75 115·1		135.1	24.5	149 149	194	73.1	2				87.1							1.101
AMIDES	Formula.	$\begin{array}{c} \text{CH}_3\text{-CONH}_2\\ \text{CH}_3\text{-CSNH}_2\\ \text{CH}_3(\text{C}_4\text{H}_3)\text{-CONH}_2 \end{array}$		$\mathrm{CH_3}(C_6H_5).\mathrm{CONH_2}$	-HNO2 H2 (0-H-2)	CH3CONH (C,H,CH2)	NO2.C6H4.CH2.NH.COCH3	C,H,CONH,					C_3H_7 . CONH.							$C_4H_9.CONH_2$
	Substance.	Acetamide Thioacetamide iso-Butylacetamide	•	Phenylacetamide	Diphenoxvacetamide	Benzylacetamide	0-Nitrobenzylacetamide	F- " Propionamide				:	n-butyramide			iso-Butyramide				Valeramide

Substance.	Formula,	M.W. (Calcd.).	Solvent.	Concentration.	M.W. (Obs.).
iso-Valeramide			Ethyl alcohol 56 Acetone 56 Chloroform 56 Benne 56	27.4-71.1 (N) 28.5-50.1 (N) 32.3-53.7 (N)	98'1-94'2 99'7-106'4 132-141
Glycollamide	CH2OH.CONH2	75	Acetone 56 Etthyl alcohol 56	11.5-16.8 (N) 32.7-65.2 (N)	73.1-81.4 77.4-79.3
Lactamide	CH ₃ CH(OH).CONH ₂	1.68	Water 57 Water 57 Ethyl alcohol 56	3.730-0.099 7.734-13.14 28.8-52.4 (N)	9.34-9.54 90.1-93.9
Urea	CO(NH2)2	9	Acetic acid 52 Ethyl alcohol 14, 38	20'9-49 6 (N) 1'48 1'32-2'64	954-111'0 67 62-63'5
Phenyl carbamide	NH2.CO.NH.C ₆ H5	136	Water 14, 1 Water 57 Ethylaicohol 56	2.815-5.371 38:2-75.8 (N)	72-74 141-159 155-164
Diphenyl carbamide	$(C_6H_5NH)_2CO$	212	Acetone 56 Acetone 56 Ethyl alcohol 56	21.7-42.4 (N) 10.0-13.0 (N) 8.0-8.7 (N)	147-100 206-220 207-196
Benzamide	C ₆ H ₅ .CONH ₂	121	Acetic acid 52	2.00	126
Thio-benzamide Acetyl benzamide	C,H,CSNH, C,H,CONH,COCH,	137	Naphthalene ¹¹ p-Dibromobenzene ⁶⁸	2.813-8.597	132-197 215-257
le iide amide aamide	C ₆ H ₄ (OH).CONH.COCH ₃ C ₆ H ₅ :COOC ₆ H ₄ :CONH ₃ C ₆ H ₅ :SO ₂ NH ₃ (CH ₃)C ₆ H ₄ :SO ₂ .NH ₃ CH.C ₂ H ₅ :CO	179 241 157 171	j-Dibromobenzene 68 p-Dibromobenzene 68 Naphthalene 58 Naphthalene 59	1.321.4'999 0'964-3'343 0'90-2'11 1'50-4'12	231-270 295-348 166-203 175-238
Ethyl succinimide	HN	127	p -Bromotoluene 9	0.980-4571	127.7-274.5

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M.W. (Obs.).	149-343 176-409 160-345 171-397 147-178 128-131 140-197 174-338 161-194	128-213 116-6-119-6 116-6-119-6 115-170 135-145 14-157 151-188 150-210 190-25 165-200 165-200 162-164	154-157 160-293 139-159 146-274 165-184 173-0-1767 145-202
Concentration.	1.26-11.20 1.37-12.87 0.67-7.69 0.54-8-41 1.28-6.16 2.17-10.10 1.90-14-42 1.90-14-42 1.50-12.05 4.501-9.026	1.13-12.08 0.96-8-65 32-4-68-(N) 23:5-4-28-(N) 18:2-22.2 (N) 0.38-0.77 1.55-10-25 1:51-797 1.79-9:33 1.58-9-62 1.64-5.01 8:04-0.94	1.74-193 1.45-9.68 1.49-13.52 1.75-13.74 1.00-8.93 1.054-4.866 1.01 8.89
Solvent.	Benzene 13, 1, 56 \$\rho \text{Nylene}^{13}\$ \$\rho \text{Chlorotoluene}^{13}\$ \$\rho \text{Drom. toluene}^{13}\$ \$\rho \text{Drom. toluene}^{13}\$ \$\rho \text{Nitrotoluene}^{13}\$ \$\rho \text{Nitrotoluene}^{13}\$ \$\rho \text{Toluidine}^{13}\$ \$\rho \text{Toluidine}^{13}\$ \$\rho \text{Dichlorbenzene}^{13}\$ \$\rho Dichlorben	Naphthalene 58 Dimethyl oxalate 24 Ethyl alcohol 56 Acetone 56 Chloroform 56 Ether 56 Ether 56 Benzene 12 Naphthalene 58	Aceic acid 38 Benzene 14, 38 Benzene 14, 38 Ethyl alcohol 14, 58, 38 Naphthalene 58 Dimethyloxalate 24 Anisol 70 Naphthalene 11 Naphthalene 11
M.W. (Calcd.).	121	137 155'5 155'5 200 166 166	151
Formula.	С ₆ Н ₅ NH.COH	C, H, NH. CSH C, H, (Cl) NH. COH C, H, (Cl) NH. COH C, H, (Br) NH. COH C, H, (NO.) NH. COH	C ₆ H ₆ NH.CS.CH ₆ C ₆ H ₄ (Cl) NH.COCH ₆
Substance.	Formanilide	Thioformanilide o-Chloroformanilide p-Chloroformanilide p-Bromoformanilide p-Nitroformanilide m-Nitroformanilide p-Nitroformanilide	Acetaninde Thioacetanilide o-Chloroacetanilide

Solvent. Concentration. M.W. (Obs.)	Naphthalene 58	H'	82 82 82 83 83 83 83 84 85 85	
M.W. (Calcd.).	169'5 169'5 214 214 180 180 180	163	149 163 163 177 331 359 151	165
Formula.	C, H, (Cl) NH.COCH, C, H, (Br) NH.COCH, C, H, (Br) NH.COCH, C, H, (Br) NH.COCH, C, H, (No.) NH.COCH,	C ₆ H ₅ N.(C ₂ H ₅).COCH ₃	C, H, NH. COC, H, C, H, NH. COC, H,	C ₆ H ₅ NH.COCH(OH).CH ₃
Substance,	m-Chlor, acetanilide p-Chlor, acetanilide m-Brom, acetanilide p-Brom, acetanilide p-Nitroacetanilide p-Nitroacetanilide p-Nitroacetanilide p-Nitroacetanilide	Ethyl acetanilide	Propioaniide Butyraniide iso-Butyraniide iso-Valeraniide Palmitinaniide Stearinaniide Glycollaniide	Lactanilide

206-220 201-269 186-185 190-196 336-300 241	185-223 207-205'5 227-226 187-189 225-257 245-291 236-323 175-231 159-197 155-194	179-395 179-395 142-266 187-257 222-298 128-374 168-374 171-309 232-330	104-212 144-197 178-258 178-204 171-268 157-270 148-308 186-200
25'8-32'1 (N) 12'8-32'5 (N) 14'8-22'6 (N) 1'19-6'75 1'49-4'08	1.79-5.83 240-39.2 (N) 1.05-3.11 30'5-43.2 (N) 1.86-8.78 1.04-4.39 0.58-5.21 1.07-8.75 2.54-10.62	0.78-10.18 0.37-7.37 0.79-4-63 0.78-5.51 0.38-13.78 1.01-6.39 0.73-5:58 0.62-4:31	1.30-5.92 1.89-11.80 1.81-10.32 1.87-9.66 1.97-3.96 2.41-13.95 1.39-9.59 1.50-9.80
Ether ⁵⁶ Benzene ⁵⁶ Ethyl alcohol ¹⁴² Aniline ³⁷ Benzoylcyclohexylamine ¹²⁸ Benzene ^{1, 142}	Naphthalene ⁸⁸ Chloroform ¹⁴² Chloroform ¹⁴² Dimethyl oxalate ²⁴ Acetone ¹⁴⁸ Naphthalene ¹¹ Naphthalene ¹¹ Naphthalene ¹¹ P.Dibrombenzene ² p-Dichlorbenzene ¹³ Methyl p-toluate ¹³ Methyl p-toluate ¹³ p-Nitrotoluene ¹³	p-todotoluene 1.9 p-todotoluene 1.3 Benzene 1.3 p-Chlorotoluene 1.3 p-Bromotoluene 1.3 p-Xylene 1.3 p-Chlorotoluene 1.3 p-Bromotoluene 1.3 p-Bromotoluene 1.3 p-Nicotoluene 1.3 p-Nicotoluene 1.3 p-Nicotoluene 1.3 p-Nicotoluene 1.3 p-Nicotoluene 1.3	Methyl p-toluate 18 Naphthalene 58
761	227 242 242 135	135	180 180 180 149 194
C ₆ H ₅ NH.COC ₆ H ₅	C,H,NH.COC,H,OCH, C,H,NH.COC,H,NO, C,H,NH.COC,H,NO, C,H, (CH,) NH.COH	C ₆ H ₄ (CH ₃) NH.COH	C ₆ H ₃ (CH ₃) (NO ₂) NH.COH C ₆ H ₃ (CH ₃) (NO ₂) NH.COH C ₆ H ₃ (CH ₃) (NO ₂) NH.COH C ₆ H ₄ (CH ₃) NH.COCH ₃ C ₆ H ₃ (CH ₃) NH.COCH ₃
Benzanilide	o-Methoxybenzanilide o-Nitrobenzanilide m-Nitrobenzanilide Formo o-toluide	Formo-\$-toluide	3-Nitro 4-Formotoluide 2-Nitro 4-Formotoluide 3-Nitro 6-Formotoluide 0-Acettoluide m-Acettoluide P-Acettoluide 2-Nitro 4-Acettoluide

142							IV.	ιυ	L.	E (٦.		A	K	1	15	5	U		A	17	OI	<i>V</i>							
M.W. (Obs.)	247-299	246-248	307-300	277-300	331-354	180 201	165-273	142-173	210-234	192-261	172-261	180-250	188-253	164-194	176-358	222-233	225-251	238-276	185-263	171-300	188-321		M.W. (Obs.),	45.7-468	57.1-59.3	56.5-63.0	145-153	84.8 88.2	88.9-ro4.r	110.4-122.9
Concentration.	0.69-2.75	1.03-6.14	3.16-5.28	20.6-60.1	1.23-3.07	0.32-0.99	1.58-18.02	0.18-0.36	0.44-0.86	0.49-1.75	0.34-1.66	1.20-12.06	1.85-12.54	1.78-13.22	1.61-17.14	L1.9-60.1	1.89-8.78	1.00-9.30	29.9-62.I	0.40-4.81	1.72-11.81		Concentration.	3.915-7.473	r.295-5.360	2.329-5.147	0.897-1.813	0.621-7.114	1.040-0.140	2.752-9.324 1.326-5:511
Solvent,	Naphthalene ¹¹ Naphthalene ¹¹	Naphthalene 11	Naphthalene 11	Naphthalene	Fyridine 3	Benzene 12	Naphthalene 58	Benzene 1	p-Chlorotoluene 13	p-Bromtoluene 18	p-Iodotoluene 13	p-Nitrotoluene 13	Methyl- p -toluate ¹³	Naphthalene 58	Naphthalene 58	Naphthalene 11	Naphthalene 11	Naphthalene 11	Naphthalene 11	Benzene 12	Naphthalene 58		Solvent,	Water 17	Benzene 17	Water 17	Formamide 42	Water 17	Benzene 1	Benzene '' Water 71
M.W. (Calcd.)	256 256			271	·		149		151					165	165	227	227	243	177	165	179	ss.	M.W. (Calcd.).	45.07		59.I	129.2	87.1		1.201
Formula.	C ₆ H ₄ (CH ₃) NH.CO.C ₆ H ₄ (NO ₃) C ₆ H ₇ (CH ₅) NH.CO C ₆ H ₇ (NO ₅)	CeH, (NO2) NH.CO (CeHs)	Cert (NO2) NH.CO CeH, (NO2)	Cens. NH. CS. Cens. (CHs.) (CC. Hs.)		CH3)2 Cansan CHO	C,H, (CH,), NH,CO CH,	4/0	C ₆ H ₄ (OCH ₃) NH.CO H					C,H, (OCH,) NH.CO CH,	Cent (OCH3) NH.CO CH3	C,H,CONH.C,H, (OCH,)	CeH5.CO NH. CeH4 (OCH3)	C,H,NH.CSC,H,OCH,	C ₆ H ₂ (CH ₃) ₃ NH.COCH ₃	C2H5OC6H4N: CHOH	C2H5OC6H4NH.COCH3	AMINES	Formula.	$C_2H_5NH_2$		$C_3H_7NH_3$	$(C_4H_9)_2NH$	$C_5H_{11}NH_2$	1111 110 11 0	C ₆ H ₅ CH ₂ NH ₂
Substance.	o-Nitrobenzo-o-toluide m-Nitrobenzo-o-toluide	Benzo-o-nitrotoluide	o-Nitrobenzo nitrotoluide	Transit di toluithioanille	a Formowylide	h-Formoxylide	m-Acetoxylide		Formo-p-anisidide					o-Acetanisidide	p-Acetanisidide	Benzo-o-anisidide	Benzo-p-anisidide	Anisithioanilide	Acet-pseudocumide	p-Formyl phenetidin	p-Acet-phenetidide		Substance.	Ethylamine		Propylamine	Di-150-butylamine	1so-amylamine	Donner	Denzylamine

Aniline	$C_6H_6NH_2$	93.1	Water 17 Benzene 17 Nitrobenzene 5 Ether 18 Naphthalene 58 Cyclohexane 21	1.976-3'891 1.828-8'828 0'5723-21'4049 1.826-2'153 1'20-6'59	10. 2-107.4 104.5-121.5 80.5-121 93.5-93.4 94-102 106-136
m-Nitro aniline	$(NO_2)C_6H_4NH_2$	138	b-Bromotoluene 9 Bromoform 4 Water 17 Benzene 17	0°338-46'011 0°2021-5'4193 1°735-2°153 2°617-7°243	86.7-226.1 87.3-94.1 174-174 140-164
 ρ-Nitroso methyl aniline ρ-Nitroso ethyl aniline ρ-Nitroso dimethyl aniline ρ-Nitroso diethyl aniline piphenylamine 	(NO) C ₆ H ₄ NH.CH ₃ (NO) C ₆ H ₄ NH.C ₆ H ₆ (NO) C ₆ H ₄ N(CH ₃₎₂ (NO) C ₆ H ₄ N(C ₇ H ₈₎₂ (C ₆ H ₅) ₂ NH	136 150 150 178 169.1	Naphthalene 24 Naphthalene 24 Naphthalene 24 Naphthalene 24 Naphthalene 24 Ethyl alcohol 17 Heesne 33	1.19-6.88 1.19-6.88 1.24-7.91 1.08-7.57 1.53-11.40 8.678-13.62	139-100 152-197 159-206 151-177 183-222 190-206
			Stearic acid 29 Benzonitrile 129	2.359-28.03 0.63-1.17	175.6-205.5 233-228
p-Nitroso diphenylamine p-Toluidine p-Anisidine a-Nitroso p-Naphthylamine Benzoylcyclohexylamine Pytrole	C, H, NH. C, H, (NO) C, H, (CH ₃) NH ₃ C, H, (OCH ₃) NH ₃ (NO) C ₀ , H, NH ₂ C, H, CONH. C, H ₃ C, H, NH	198 107 123 172 203 67	Naphthalene 34 Naphthalene 58 Naphthalene 58 Naphthalene 24 Benzamilide 128 Cyclohexane 21 Naphthalene 24 Senzamilide 128 Cyclohexane 21 Naphthalene 21	1.25.8:57 1.25.8:57 1.05-7:94 1.23.11:27 0.70-5.14 1.91-4:49 0.64-2:07 0.368-10:23	203-251 95-112 111 140 180-227 184-194 84-113 89-109
N-Methyl pyrrole N-Ethyl pyrrole	$C_4H_4N.(CH_3)$ $C_4H_4N.(C_9H_1)$	81 95	Acetic acid we Benzene 50 Acetic acid 50 Benzene 50	0.245-7.006 0.245-7.006 0.840-27.65 0.361-770	79-85 81-111 87-94
a-Acetyl pyrrole a-6-Dimethyl pyrrole Tetra-iodo pyrrole	CH ₃ .CO.C ₄ H ₂ NH (CH ₃) ₂ .C ₄ H ₂ .NH C ₄ I ₄ NH	109 95 571	Acetic acid ⁵⁰ Benzene ⁵⁰ Acetic acid ⁵⁰ Benzene ⁵⁰ Acetic acid ⁵⁰ Acetic acid ⁵⁰	0.248-29'86 0.229-79'56 0.333-17'72 0.472-16'83 0'368-14'52 0'404	96-132 111-176 112-121 90-127 98-119 523

M.W. (Obs.).	545-719 109-114 110'2-147'9	220'2-57'7 87'4-100'7 93-100 73'66-87'0	113-143 108-122 237-264		468.7-514.4		M.W. (Obs.).	72-80 91-109 141-149	163-227	232.2-246	232.0-240.4 180.4
Concentration.	0.069-2.430 0.80-2.93 0.397-30.52	0.535-4°30 2°485-17°35 0°91-5°16 4°385-2°755	3.4977-24.06 0.55-7'67 0'28-1'23		0,21-0,31		Concentration. M.W. (Obs.).	0.28-2'30 0'24-4'79 2'34-6'43	0.30-5.82	1.24-2.63	1.64-3.26
Solvent.	Ethylene bromide 50 Cyclohexane 21 ρ -Bromotoluene 9	Nitrobenzene ⁵ Naphthalene ¹¹ Water ⁷²	Nitrobenzene ⁵ Benzene ¹² Benzene ^{1, 12}		$ ho$ -Toluidine 73		Solvent.	Benzene ¹ Benzene ¹ Benzene ¹	Denzene ¹ Benzene ¹ Benzene ¹³²	Benzene 132	Benzene 132 Acetone 132
M.W. (Calcd.).	79	85	113 188		262	NITROSO COMPOUNDS.	M.W. (Calcd.).	74 102 150	178	135	135 149
Formula.	C_6H_5N	$C_{\mathfrak{s}}H_{\mathfrak{U}}N$	$\begin{matrix} C_{\rm c}H_{10}{\rm NCOH} \\ CH_3 & C & CH \end{matrix}$	CH ₃ . N CO N. C ₆ H ₅ NH NH	C ₆ H ₄ C:C C ₆ H ₄	NITROSO C	Formula.	(CH ₃) ₂ N.NO (C ₂ H ₃) ₂ N.NO (C ₄ H ₃) ₂ N.NO (C ₄ H ₃) ₂ N.O	$(NO) C_6 H_4^{1N} (C_9 H_5)_2 (NO) C_6 H_4^{1N} (C_9 H_5)_2$	(NO) CeH (CH3)	$(NO) C_6H_2 (CH_3)_3$
Substance.	Pyridine	Piperidine	Formyl-piperidine Antipyrine		Indigo		Substance.	Dimethylnitrosoamine Diethylnitrosoamine Nitrosoethylaniline Nitrosodimethylaniline	Nitrosodiethylaniline Nitroso 2'5-xylene	Nitroso 2.4-xylene Nitroso 2:6-xylene	Nitroso 2'4'6-mesitylene

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Substance.	Formula.	M.W. (Calcd.).	Solvent.	Concentration.	M.W. (Obs.).	
Urethane	$\mathrm{NH_2.CO.OC_2H_5}$	1.68	Naphthalene 11 Water 57 Benzene 56 Etthyl alcohol 56 Stearic acid 29 Acetone 56 Chloroform 56 Fthar 56	0.98-9.41 2.084-5.338 30-2-58-4 (N) 35.3-89-4 (N) 1.066-23.26 26:5-49.2 (N) 31-2-54.1 (N)	98-167 91-88-4 129-140 94-6-91*1 99'74-125'3 92'0-95'6 106'9-1121	
Phenylurethane	C ₆ H ₅ NH.CO.OC ₂ H ₅	165	Ethyl alcohol 56 Acetone 56 Chloroform 56 Ether 56 Benzene 56,1	25.2-50.0 (17) 267-55-6 (N) 247-5-11.4 (N) 2172-32.7 (N) 237-45-1 (N) 239-693.8 (N)	102-104 162-160 171-5-179 169-172	211 1 11
Methyl carbamate Phenylxanthogenamide	NH2CO.OCH3 C6H6N.C (SH).OC2H6	75 181	Naphinaene Benzene ¹ Benzene ¹	0.28-1.05 0.35-7.35	1/5-194 83-102 154-216	2122
	HIIN	NITRILES.				
Substance.	Formula.	M.W. (Calcd.).	Solvent.	Concentration.	M.W. (Obs.).	
Valeryl acetonitrile Salicylnitrile 5-Bromosalicylnitrile 3:5-Dibromosalicylnitrile 6-Nitrosalicylnitrile 3:5-Dinitrosalicylnitrile 3:5-Dinitrosalicylnitrile 3-Nitro 5-bromosalicylnitrile Salicylnitrilebenzyl ether	(CH ₃) ₃ . C.CO.CH ₃ CN C ₄ H ₃ (OH) CN C ₆ H ₃ (OH) Br.CN C ₆ H ₃ (OH) Br.CN C ₆ H ₃ (OH) NO ₂) ₂ CN C ₆ H ₃ (OH) (NO ₂) ₂ CN C ₆ H ₃ (OH) Br (NO ₂) ₂ CN C ₆ H ₃ (OH) Br (NO ₂) ₂ CN C ₇ H ₂ (OH) Br (NO ₂) ₂ CN	125 119 198 277 164 209 243 209	Benzene ¹³⁷ Naphthalene ²⁰ Naphthalene ⁷⁴	0.62-6'89 0'50-8'02 0'437-2'734 0'433-2'913 0'278 0'458-2'858 0'40-2'808	124'1-175'3 125-251 195-228 296-288 181 202-207 249-242 212-217	_

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M.W. (Obs.).	57.2-71.0 57.6-63.7 56.0-61.8	78.2-88.5	103-101	85.2-93.3	123-116 130	115-136	124-150	109.3-246.0	147-200	163-226	203-327	344-541	189	195.8-217.8	182-187	191-170	209-240	234-244	232	246-272	245-326		106'8-199'8 454'5-560'3 151'4-153'7 98'0-102'1
Concentration.	0.475-13'98 0.502-15'91 6'62-18'52	0.84-2.84	0.25-6.95	0.62-56.0	0.41-3.65 1.86	0.957-30.17	0.41-10.72	69.8-66.0	2.10-8.72	10.8-68.1	18-8-14-1	1.54-7.69	2.20	0.47-5.35	1.62-3.51	0.00-2.50	1.86-7.85	0.10-0.49	11.0	1.87-7.93	2.06-8-67		1.07-7.47 0.364-6.28 1.21-6.25 0.69-5.97
Solvent.	Benzene 8 Acetic acid 8 Water 8	Cyclohexane 127 Dicyclohexane 124	Benzene 1	Cyclohexane 127	Antimony trichloride 75 Acetic acid 52	Benzene 8	Acetic acid 5 Cyclohexane 127	Dicyclohexyl 124	Naphthalene 24	Naphthalene 24	Naphthalene 24	Naphthalene 24	Ether 18	Cyclohexane 127	Carbon disulphide 14	Antimony trichloride	Naphthalene 24	Benzene 1	Benzene 1	Naphthalene 24	Naphthalene 24		Dicyclohexyl ¹²⁴ Cyclohexane (cryos.) ¹²⁷ Cyclohexane (ebul.) ¹²⁷ Benzene (cryos.) ¹²⁷
M.W. (Calcd.).	28		100	72	120				136	150	178	332	182			,	198	214	230	243	243		86
Formula,	(CH ₃) ₂ CO		CH3.CO.CH2.CO.CH2	CH, CO.C, H,	CH3.CO.C6H5				C,H, (OH) CO.CH,	C,H, (OH) CO.C,H,	C,H, (OH) CO.C,H,	C ₆ H ₄ (OH) CO.C ₁₆ H ₃₁	C,H,CO.C,H,				C ₆ H ₄ (OH) CO.C ₆ H ₅	C ₆ H ₃ (OH) ₂ CO.C ₆ H ₅	C,H, (OH), CO.C,H,	C ₆ H ₄ (OH) CO.C ₆ H ₄ (NO ₂)	C ₆ H ₄ (OH) CO.C ₆ H ₄ (NO ₂)		$C_6H_{10}O$
Substance.	Acetone		Acetyl acetone	Methyl ethyl ketone	Acetophenone				b-Hydroxyacetophenone	b-Hydroxypropiophenone	b-Hydroxyvalerophenone	p-Hydroxypalmitophenone	Benzophenone				p-Hydroxybenzophenone	Dihydroxybenzophenone	Trihydroxybenzophenone	m-Nitro p-hydroxybenzophe-	none p -Nitro p -hydroxybenzophe-	none	Cyclohexanone

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97.1-103.6 99:0-107.8 175:4-212.5 155:0-191:6		M.W. (Obs.).	34.4 149-163 179-162	139-144 181-184 109-151	151-160 178-259 139-102	98°5-100°4 102-114 109-113	81.7-135.8 105.0-152.0 130-142 121-131	112-125 114-154 160-271 131-241	122-221 147-177 133-162	142-202 130-186 141-201
0.82-4.34 0.76-6.62 0.53-9.33 0.69-10-85		Concentration.	2.742 0.739-32:25 0.763-29:92	0.887-1519 0.372-0.719 0.267-34.07	1.113-10-69 0.568-42.12 0.574-36-37	1.858-21.70 0.511-17.64 0.452-2629	0.589-24.24 0.85-4.02 0.72-8.80 I.07-5.78	1.00-16.4 1.00-16.40 1.01-4.68 0.33-2.53 0.58-2.04	0.41-10.30 0.63-2.17 0.99-5.79	0.95-8.01 0.93-7.63 0.92-6.75
Benzene (ebul.) 127 p-Xylene 127 Cyclohexane 137 Cyclohexane 127		Solvent.	Water 76 Benzene 8 Acetic acid 8	Water 8 Benzene 8 Acetic acid 8	Water 8 Benzene 8 Acetic acid 8	Water 8 Benzene 8 Acetic acid 8	Nitrobenzene ⁵ Dicyclohexyl ¹³⁴ Nitrobenzene ²² m-Dinitrobenzene ²² 6 Diltromohan ²²	Benzil 22 Naphthalene 23, 23, 19 p -Dibromobenzene 22 p -Chlorobromobenzene 23 p -Chlorobromobenzene 23 p -Chlorobromobenzene 23	Naphthalene 19, 24, 23, 22 Nitrobenzene 22 m-Dinitrobenzene 22	\$\theta\text{Nitrotoluene}^{22}\$ 2.4-Dinitrotoluene \(^{22}\) 2.4-6-Trinitrotoluene \(^{23}\)
112	DES.	M.W. (Calcd.).	30 147.5	165.2	193.5	106	122	122	122	
$C_{7}H_{13}O \\ C_{7}H_{12}O$	ALDEHYDES	Formula.	н.сно ссі _з .сно	CC1 ₃ .CH (OH) ₃	CCI3.CH (OH) OC2H5	C ₆ H ₅ .CHO	С ₆ Н ₄ (ОН).СНО	С, н, но. сно	$\mathtt{C_6H_4HO.CHO}$	
$o ext{-}Methylcyclohexanone} p ext{-}Methylcyclohexanone}$		Substance.	Formaldehyde Chloral	Chloral hydrate	Chloral alcoholate	O * Benzaldehyde	Salicylaldehyde	m-Hydroxybenzaldehyde	$ extit{$ extit{ℓ-Hydroxybenzaldehyde}}$	

Substance.		3-Nitro-p-hydroxybenzaldehyde	3-bromo-p-nyaroxybenzalde- hyde	hyde	hyde	3.5-Ditatonyuloxybenzaute- hyde Acetyl-o-Amidobenzaldehyde Acetyl-p-Amidobenzaldehyde o-Homosalityyaldehyde	o-Homo-p-Hydroxybenzalde- hyde Homo f hydrombenzalde	m-tronto-p-nymoxyocuzatue: Benzoylaldehyde p-Thymotinaldehyde Vanillin 4-Hydroxy-a-Naphthaldehyde		Substances.	Benzene
Formula.		С ₆ Н ₃ (ОН) NO ₂ .СНО	C ₆ H ₃ (OH) Br.CHO	С ₆ Н ₂ (ОН) Сі ₂ .СНО	C_6H_2 (OH) Br_2 .CHO	C,H,CHO NI.CHO C,H,CHO NH.COCH; C,H,CHO NH.COCH; C,H,CH3 (OH).CHO	С, Н, СН, (ОН). СНО	C,H,CH, (OH).CHO C,H,CO.CH,CHO C,H, (OH).CH,CHO C,H, (OH).CH,CHO C,H, (OH) OCH,CHO C,H, (OH) OCH,CHO	HYDROCARBONS, NITRO AND OTHER DERIVATIVES.	Formula.	$C_{\mathfrak{g}}H_{\mathfrak{g}}$
M.W. (Calcd.).		167	201	161	280	374 163 163 136	136	136 148 178 152	AND OTHE	M.W. (Calcd.).	78
Solvent.	\$\rho\$-Chloronitrobenzene \$^{22}\$ \$\rho\$-Dichlorobenzene \$^{22}\$ \$\rho\$-Dibromobenzene \$^{22}\$ \$\rho\$-Chloromobenzene \$^{22}\$ \$\rho\$-Chloromobenzene \$^{22}\$ \$Naphthalene \$^{24}\$, \$^{25}\$ \$Naphthalene \$^{24}\$, \$^{25}\$	p-Azoxyanisole ²⁴ Naphthalene ²⁵	Naphthalene 25	Naphthalene 25	Naphthalene 25	Naphthalene 25 Naphthalene 58 Naphthalene 58 Naphthalene 19	Naphthalene 24, 19	Naphthalene ²⁴ Benzene ¹² Naphthalene ¹⁹ , ²⁴ Naphthalene ²⁴ , ¹⁹ Naphthalene ²⁴	R DERIVATIVES.	Solvent.	Acetic acid ^{10, 8} Phenol ¹⁰
Concentration.	1.00-6'24 0'26-0'71 1'03-7'75 0'96-9'65 0'31-0'91	0.344-0.998 1.11-8'95	1.76-10-22	1.05-939	1.02-8.50	1.08-9.91 1.63-8.77 1.63-3.50 0.43-5.02	0.47-3.53	0°96-7°17 0°54-0°95 0°66-9°92 0°59-10°04 0°80-2°65		Concentration.	0.89-25.47
M.W. (Obs.).	150-196 152-201 207-604 135-154 150-254 159-317	125-124 164-176	210-259	187-220	266-307	373-418 160-173 166-202 139-140	133-191	154-255 101-115 190-267 147-185 179-234		M.W. (Obs.).	79.6-113.1

126-143 148-155	144-143 171-159 2010-2051 64'9-72 70'8-101'4	57-03 182-6-208-7 911-135-7 75-2-76-6	84-94 131-143 126-166	119-149 181-211 170-175 202-246 82-3-93-2 85-2-100-8	78'0-105'4 164-195 179-186 432-346	M.W. (Obs.).	47.4-49.3	122-124 129 170	117.5-131'1 247-314 262'3-279'4
0.455-6.008	2.81-9.02 1.96-7'44 1.261-3'108 0'681-2'718	0.29-2:32 1.27-4.50 0.37-3:37 0.844-7:56	0.27-4.18 2.555-5.891 0.779-29'68	0.902-32-32 5.912-12-57 2-733-4-661 1.100-3-567 0.291-5-48 0.519-19-55	0'498-22'b0 0'3096-3'346 3'187-5'332 1'257-3'718	Concentration.	0.127-0.342	0.23-0.60 1.581-18:86	0.435-1.583 0.435-1.583 0.1572-21.00 2.423-7.197
Acetic acid 8 Ethyl alcohol 14	Acetonitrile ¹²⁹ Acetonitrile ¹²⁹ Anisol ⁷⁰ Nitrogen peroxide ¹²⁴ Cyclohexane ¹²⁵	Benzene ¹ Dicyclohexane ¹²⁴ Cyclohexane ¹²⁵ Nitrogen neroxide ¹³⁴	Benzene 1 Ethylene bromide 62 Acetic acid 8, 52	Denzene Ethyl alcohol ¹⁷ Hexane ¹³³ Hexane ¹³³ Bromoform ⁴ Nitrobenzene ⁵	p -Bromotouene 9 Chloroform 27 Benzene 27 Acetone 114	Solvent.	Iodine (cryos.) 77 Lodine (ebul) 78	Methylene iodide 49 Carbon disulphide 79 Carbon disulphide 79	Sulphur (ebul.) 83 0435-17583 Carbon disulphide ^{79, 80, 88, 90} 1-572-21 00 Naphthalene ^{80, 90} 2-423-7'197
128	154 178 61	.194 75	89 123	168 153 183 84	357	ELEMENTS. M.W. (Calcd.).	54	123	123 c 75°0 256
$C_{10}H_8$	$\begin{array}{ccc} C_{12}H_{10} & & & \\ C_{14}H_{10} & & & \\ CH_{3}NO_{2} & & & \end{array}$	$\begin{array}{c} C \ (NO_2)_4 \\ C_2H_5NO_2 \end{array}$	$(CH_3)_2$ $CHNO_2$ $C_6H_5NO_2$	C ₆ H ₄ (NO ₂) ₄ (NO ₂) C ₆ H ₄ OCH ₃ NO ₂ C ₆ H ₃ ; (OCH ₃) ₂	${ m C_{10}H_8.C_6H_2~(NO_2)_3OH} \ { m C_{10}H_{16}O_3N_2}$	ELEM Formula.	Al_2	${ m P_4}$	As S _s
Naphthalene	Diphenyl Anthracene Nitromethane	Tetranitromethane Nitroethane	secNitropropane Nitrobenzene	 m-Dinitrobenzene ρ-Nitroanisol Dimethoxynitrobenzene Thiophene 	Naphthalene picrate Bornylene nitrosite	Substance.	Aluminium	Phosphorus	Arsenic Sulphür

M.W. (Obs.).	292-242 254:5-257:9 249-250 246-249 52:5-62:8 217-274 43:6-89:0 337-351 1899-195*4 234:6-249*4	254 225-263 239-260	204-270 238-251 233-256 624-614	191.6-191'3 796 64-69 169'4-169'1 159-171'2	II2'5-IIO M.W. (Obs.).	1481-612 1018-629 63.7 765-686 688-756 63-696 82-6-896
Concentration. M.W. (Obs.).	0.79-2.49 0.3094-1.048 0.728-1.48 0.73-1.05 0.73-0.93 0.92-8.58 0.3186-0.3279 0.2599-0.8501 0.33-2001	? 0.38-2.36 0.38-2.76	0.54-4.19 C.07-2.57 0.78-3.76 0.203 0.480	0'905-4'066 0'302 0'94-2'93 I'19-3'55 0'34-1'86	o'os47-o'311 Concentration.	0.291-2.432 0.036-1.744 } 0.87-2:97 0.24-2:23 0.96-3:83
Solvent.	Fenchone ⁸¹ Bromoform ⁸² Aluminium bromide ⁸⁴ (cryos.) Methylene iodide ⁴⁹ Bromine ⁸⁵ Chlorine ⁸⁵ Chisulphur dichloride ⁸⁵ P. Nitrotoluene ⁸⁶ Benzene ⁸⁷ Phosphorus ⁸⁹ Toluene ⁸⁹	Diphenyl 90 Iodine 102 Tin tetrachloride 85	Arsenc trichloride 52 Tin tetrachloride 85 Arsenic trichloride 85 Phosphorus 91	Sulphur ** (ebul.) Methylene iodide 49 Disulphur dichloride 85 Sulphur ** (ebul.)	Jouine ··· (ebul.) Solvent,	Nitrobenzene 44 Benzene 44 Acetic acid 2 Nitrobenzene 92, 135 Ethylene bromide 136, 92 Acetic acid 99, 135 Nitrogen peroxide 134
M.W. (Calcd.).		256	256 256 79.2	127'5	ACIDS. M.W. (Calcd.).	36·5 63
Formula.		<i>യ്</i>	<i>ೲ</i> ຶ៷៓៷៓	T e	Formula,	HCl HNO ₃
Substance.		Sulphur (rhombic)	Sulphur (amorphous) Sulphur (monoclinic) Selenium	Tellurium	Substance.	Hydrochloric acid · Nitric acid

66.3-71.5 58.8-59.7 252.6-190 1794-209.1 182.0-142.7 0.79-100.9	67.4 52.1-59.8 55.7-65.0 137.9-143.8	M.W. (Obs.). 18·9-23·5 21·2-23·8 22·1-23·8 28·69-33·29 20·6-34·9	21.3-32.5 17-6-30.0 18'5-32'4 19'38-19'62 83'2-83.0 94-6-90'3 105-7-108'2 142-15.2 288-455 113-238 371-402 39-4402	192-179 167-193 275-290 258-253
0.93-5.40 1.78-3.19 1.74-8.42 1.95-7.66 0.35-1.10	. 0'76 . 0'76 . 0'058-1'78 2'35-14'74 0'403-19'15	Concentration. o o 105-0 o 105 o 0 144-0 o 236 o 0 305-0 o 35 o 0 305-0 o 105 o 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	0.236.213 0.0539-1.121 0.0539-1.132 0.041-1.732 0.92-8.97 0.92-8.97 0.96-1.169 0.96-2.09 0.92-1.97 0.59-3.01 0.59-3.01	0.526-2.59 1.261-5'99 0'852-4'32 0'407-3'67
Monochloroacetic acid ¹³⁵ , ⁹² Phosphorus oxychloride ²⁸⁶ Acetic acid ¹³⁵ , ² Nitrobenzene ¹³⁵ Monochloroacetic acid ¹³⁵ Benzene ⁹⁹	Phosphorus oxychloride 126 Phosphorus oxychloride 126 Formamide 42 Phenol 87 Formamide 42	Solvent. Bromoform ⁷ Ethylene bromide ⁷ Dimethylaniline ⁷ p-Toluidine ²⁹	Menyl oxalate ' Methyl succinate 7 Veratrol 7 Veratrol 7 Acetic acid 10 Phenol 13, 10 Benzene 138 Acetic acid 139 Phosphorus oxychloride 131 Benzene 130 Sulphuric acid 34 Sulphuric acid 34 Sulphuric acid 34 Antimony trichloride 35 Fenchone 31 Vitachloride 31	Sulphuric acid ⁹⁴ Sulphuric acid ⁹⁴ Sulphuric acid ⁹⁴ Sulphuric acid ⁹⁴ Phosphorus oxychloride ¹⁸¹
86	82 62 138	OXIDES. (Calcd.). 18	76 92 108 220 140 198	230 . 182'9 144 255
H ₂ SO ₄	H ₃ PO ₃ B (OH) ₃ (CH ₃) ₃ ASOOH	Formula. $ m H_2O$	N N N N N N N N N N N N N N N N N N N	As,O; C1,O7 MoO3 OsO4
Sulphuric acid	Phosphorous acid Boric acid Cacodylic acid	Substance. Water	Nitrogen trioxide Nitrogen peroxide Nitrogen pentoxide Phosphorous oxide Boric anhydride Arsenious oxide	Arsenic oxide Chlorine heptoxide Molybdenum trioxide Osmium tetroxide

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S. Carteria	COMPOUNDS INDICATED THOUGHNIC COMPOUNDS	ORGAINIC CO	MPOUNDS.		
ounstance.	Formula.	M.W. (Calcd.)	Solvent.	Concentration.	M.W. (Obe.)
Phosphorus trichloride Phosphorus oxychloride	PCI ₃	136.33	Nitrobenzene 130	0.63-1.35	124.6-125.8
Hydrogen phosphide (solid)	$_{\mathrm{HP}_{2}}^{\mathrm{LG}_{3}}$	153°35 63	Monochloroacetic acid 131 Phosphorus 93	0.29-5.47	125.8-179.9
Disulphur dichloride	SCI.	103	Sulphur dioxide 39	0.52-4.82	300-375 216-230
Iodine monochloride	2 - 2	134 9	Sulphur dioxide 35 Bromine 85	1.99-5.53	140-146
	7	162.4	Phosphorus oxychloride 151	2.06-5.88	168-164
a-todine monochloride 8-Iodine monochloride	IG		Bromoform 151 Bromoform 151	0.611-7.56	164-165 166-167
rodine cyanide	ICN	153	Phosphorus oxychloride 151	0.015-0.954	173-184 150-156
•	S	SALTS.			
Substance,	Formula,	M.W. (Calcd.).	Solvent.	Concentration.	M.W. (Obs.).
Lithium chloride	LiCI	42.5	n-Propyl alcohol 94	0.1824-5.01	36.10-48.05
			Bismuth chloride 95 Ethyl clockel 97 14 38	0.290-0.552	45'3-44'5
			iso-Amyl alcohol 96	0.658-6.060	34.03-29'16
			Formic acid 41	0.842-1.885	26.53-25.07
			iso-Butyl alcohol 97	2.01-6.06	09.9-07.74
Lithium bromide	LiBr	86.86	iso-Amyl alcohol 97 Ethyl alcohol 97	0.41-5.55	41.79-44.42
			iso-Butyl alcohol 97	1.36-6.38	02'01-57'73 83'51-00'53
Lithium iodide	Lil	133.86	iso-Amyl alcohol 97 Fthyl alcohol 97	1.28-6.41	89.82.87.24
		66	iso-Butyl alcohol 97	5.70-11.02	120.8-141.7
			iso-Amyl alcohol 97	96.11-22.1	154.4-157.5
Tithing nitrate			Acetic acid " Iodine 77 (crvos.)	1.21-2.13	192.5-230.9
Trainin muale	Lino	68.89	Ethyl alcohol 97,	2.82 5.65	53.83.56.65
			150-Butyl alcohol 97	0.50-33.1	74.57-82.26
Lithium acetate	CH,COOL;	66.03	Acetic acid 97		78.05-102.3
	,			0.059-2.073	02-00

Lithium salicylate Sodium chloride	C ₆ H ₄ (OH) COOLi NaCl	144 58.5	n-Propyl alcohol 94 Bismuth chloride 95 Macouric chloride 100	1.973-15.45	246.8-288 9 65 0-64.2
			Mercuric chloride	0.51-2.00	36.85-38.50
Sodium bromide	NaBr	103	Formic acid 47	1.522-4.995	76.22-72.10
Sodium iodide	NaI	150	Acetic acid ** Ethyl alcohol 14	0.56-14.35	73.11-82.33
		,	Sulphur dioxide 69	39'53	265
			n-Propyl alcohol 94	3.09-14.02	140.1-150.5
			Iodine 77 (cryos.)	0.109-0.268	445-878
S. dina	00		Acetic acid %	0.29-1.51	102.1-166.7
Sodium sulphate	Na ₂ SO ₄	142	Sulphuric acid 34	0.71.2.08	122-100
Sodium pyrosulphate	Nas Co	222	Sulphuric acid	1.27-2.39	234-250
Souluin acetate	CH3COONA	82	Acetic acid 12, 33	16.0-11.1	1.22-0.60
Potassium chloride	KCI	4.6	Filenol Formic acid 41	0.1015-3.010	40.0-62.9
		٠+ ٢	Bismuth chloride 95	0.487-12 06	70.8-72.4
			Mercuric chloride 100	0.11-0.50	77.3 75.0
Potassium bromide	KBr	0.611	Formic acid 41	2.201-6.542	80.9-77.3
Potassium iodide	KI	166	Sulphur dioxide 39, 69, 66	1.85.14.58	235-317
			Iodine 77 (cryos.)	0.158-0.891	293-265
			Iodine 102, 78 (ebul.)	0.78-1.75	229.6-240.5
			Acetamide 140	0.212-2.226	81.2-82.6
Potassium sulphate	K_2SO_4	174	Sulphuric acid 94	1.74-3.17	136-130
Potassium hydrogen sulphate	KHSO,	136	Sulphuric acid 94	2.58-2.67	229-245
Potassium pyrosulphate	$K_2S_2O_7$	254	Sulphuric acid 94	1.91-4.76	
Potassium thiocyanate	KCNS	26	Sulphur dioxide 69, 66	146.7-454.5(N)	_
Potassium acetate	CH3COOK	98.15	Acetic acid 99	0.979-3.214	93-92
			Ethyl alcohol 14, 38	1.16-10.87	82-102
Rubidium chloride	RbCi	120.6	Bismuth chloride 95	0.830 - 1.267	9.511-2.611
			Mercuric chloride 100	0.28-1.16	211-701
Rubidium iodide	RbI	212	Sulphur dioxide 69, 66	72.46 (N)	376
			Jodine 77 (cryos.)	0.307-1.236	303-314
	į	•	lodine 'o (ebul.)	0.307-1.80	244-307
Caesium chloride	CsCl	168.4	Bismuth chloride 50	0.822-1.743	169.5-177.2
J. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	H	1	Mercuric chloride 100	0.34-1.13	153-158
Caesium logide	CSI	2599	Lodine '' (cryos.)	0.400-1.012	380-373

154			MUL	LCOLAR .	1550017	111014		
M.W. (Obs.).	251-334 45'9-53'8	74.07-55.82 345 227	102'5-106'0 125-155 107-174 118'7-122'1	121-150 121-166 125-8-127-2 136-06-139-41 130-0-147-1	70.5-194 9 271.1-263.7 76.5-81.3 148'90-145'39 147-132	221-219 175'30-184'5 328'29-360'08 140'8-145'99	438:15-442'56 462'8:480'83 173'6 188'32-383'83	220-239 135-144 237-228 207-184 78'46-84'50
Concentration.	0.268-1.490 0.07-0.34	1.769-3.893 93.46 (N) 7.95 (N)	0390-0977 032-077 0289-1093 041-082	0.45.415 0.583-0.983 1.58-4.02 5.35-4.23 0.78-2.508	0.48-0.77 0.801-2.127 0.49-3.14 1.60-5.18	1.50 5.39 0.14-1.08 0.82-2.47 1.046-2.189	0.94-4.13 0.82-2.18 0.37 0.97-609 1 65-17-68	149-3.51 1.69-9.42 0.586-1.911 0.290-1.749 0.383-0.763
Solvent.	Iodine ⁷⁸ (ebul.) Mercuric chloride ¹⁰⁰ Formic acid ⁴¹	Formic acid 41 Sulphur dioxide 69, 66 Sulphur dioxide 66	Bismuth chloride 95 Mercuric chloride 100 Quinoline 103 Pyridine 104 Methal cultified 104	Ethyl sulphide 104 Bismuth c. loride 95 Pyridine 104 Methyl sulphide 104 Ethyl slochol 95 Ethyl alcohol 95 A come 98	Acctanical 136, 140 Acctamide 140 Pyridine 104 Methyl sulphide 104	Etnyl sulpnide 107 Pyridine 104 Piperidine 104 Bismuth chloride 95	Piperidine ¹⁰⁴ Piperidine ¹⁰⁴ Piperidine ¹⁰⁸ Urethane ¹⁰⁸ Piperidine ¹⁰⁹ Pyridine ¹⁰⁸	Denzontrile ¹²⁹ Acetonitrile ¹²⁹ Pyridine ⁷² Pyridine ⁷² Pyridine ¹⁰⁶
M.W. (Calcd.).	53.4	97'9 145 76'2	99.05	134'5 134	142'99	142'9 89'16 286'06 143'4	374.84 468.40 170 169.61	134 166 79.77
Formula.	NH,CI	NH, Br NH, I NH, CNS	Cuci	CuCl,	CuBr	CuCN AgCI	$egin{array}{l} AgBr \\ AgNO_3 \end{array}$	Agen Agens Giel
Substance.	Ammonium chloride	Ammonium bromide Ammonium iodide Ammonium thiocvanate	Cuprous chloride	Cupric chloride	Cuprous bromide	Cuprous cyanide Silver chloride	Silver bromide Silver nitrate	Silver cyanide Silver thiocyanate Glucinum chloride

Mornecium formate	(HCOO) _o Mg	114	Formic acid ²	۸۰,	529.6	
Magnesium costate	(CH.COO), Mg	142	Acetic acid 2	۸.	304.3	
Magnesium accate	CaCl	III	Bismuth chloride 95	0.227-0.356	113.8-115.1	
Coloium iodide	Cal.	293.8	Iodine 102	1.81-4.67	1043-1105	
Calcium contate	(CH,COÔ),Ca	158	Acetic acid 99	c.839 - 1.824	214-226	
Carcium acetate	SrCI.	158.6	Bismuth chloride 95	0.212-0.460	163.9-165.4	
Strontium circuit		341.3	lodine 102	1.65-4.95	2201-9.006	
Strontium costate	(CH.COO).Sr	205.6	Acetic acid 99	0.839 1.824	214-226	
Strontum acctate	BaCl.	208.3	Bismuth chloride 95	612.1-122.0	210.0-214.6	
Parium conorde	(CH,COO),Ba	255.4	Acetic acid 99	1.500-4.839	253-254	
Zino oblonide	ZnCl	136.3	Bismuth chloride 95	0.608-1.037	137.9-141.5	
Zinc cirioride	N i)	Quinoline 103	0.195-6.772	146-137	
		136	Piperidine 104	1.45-2.65	147.69-168.36	
		•	Urethane 136	0.10	138.7	
			Pyridine 104	0.6-96.0	125.07-129.23	
			Methyl sulphide 104	0.93-4.18	245-211	
	ZnBr.	225.3	Ouinoline 103	0.721-3.535	223-226	
Zinc bromide	2	224.4	Pyridine 104	0.87-3.15	212.96-208.54	
	Þ	225	Methyl sulphide 104	0.03-4.18	245-211	
			Aluminium bromide 84	2.93-8.12	215 220	
	ZnI	310.1	Quinoline 103	0.498-2.381	306-325	
Zille louide	N	n D	Methyl sulphide 104	1.61-7.59	310-316	
Codming chloride	CdCl	183.3	Bismuth chloride 95	0.887-1.491	1.87.3-193.1	
Cadilliain Circuia	4)	Quinoline 103	r.300-7.498	183-187	
			Urethane 186	0.44	1.83.1	
Cadmium bromide	CdBr,	272.3	Quinoline 103	5.618-10.8 2	5203-522	
	4	271.22	Piperidine 104	2.10-6.62	280.2-329.4	
		272	Aluminium bromide 84	0.55		
Codmin indide	CdI	366.3	Quinoline 103	2.287-11.57		
Caumining round		366	Ethyl alcohol 14, 38, 108	2.59-27-84		
)	Water 14, 38, 107, 108	4.36-21.20		
			Piperidine 104	1.78-14.59	343.1-397.7	
			Pyridine 104	0.80-3.20	395.9-3-5-5	
			Methyl sulphide 104	1.32-8.48	340-372	
	1		Ethyl sulphide 104	1.73-0.52	350-375	
Mercuric chloride	$_{ m HgCl}_{ m s}$	271	Ethyl alcohol 17, 23, 23	2.03-30.00	402-604	

Substance.	Formula.	M.W. (Calcd.).	Solvent,	Concentration. M.W. (Obs.).	M.W. (Obs.).
			Water ⁵³ , 38	8.978-4.253	260.8-263.3
			Methyl sulphide 104	0.84-3.60	329-335 238-275
			Ethyl sulphide 104	2.56-10-46	272-280
			Benzonitrile 139, 104 Fenchone 81	1.72-4.02	300-335
			Urethane 136	1.12-9.07	275-282
Mercurous chloride	$ar{f H}{f g}_2{f C}{f I}_2$	471	Mercuric chloride 100	0.20-3 /0	471-450
ricically biolillag	$HgBr_2$	360	Fenchone 90	1.02-6.47	355-369
Mercuric iodide	HgI	727	Aluminium bromide 84 Ethyl alcohol 109	0.18-5.95	346-500
	•	·	Pyridine 104		283.8-318.0
			Methyl sulphide 104	1.89-12.23	463-460
			Ethyl sulphide 104		464.452
			Benzonitrile 104, 129		435-513
Mercury methyl chloride	HgCH,Cl	25.1	Fenchone 31 Methyl sulphide 104	0.92-4.95	403-458
	4	,	Ethyl sulphide 104		203-240
Mercury methyl iodide	HgCH,I	342	Methyl sulphide 104		254-259
Money	·)	-	Ethyl sulphide 104		303-320
The flame of flame	$Hg(CN)_2$	252	Pyridine 104		220.8-210.8
Aliminium chlorida		239.5	Mercuric chloride 100		218-233
	AIC13	133.5	Ether 49		137-189
			Fyridine 104 Nitrobenzene 110	0.72-2.38	125.9-128.4
			Aluminium bromide 84	0.50-4.32	125-131
Aluminium bromide	AlBr ₃	267	Bromine 49, 85	3.453-11.50	528-584
			Carbon disulphide 110	i.5-10.9	539-540
Aluminium iodide	11.4	q	Nitrobenzene 110	8.22-15.61	275-277
	AIL3	400	Aluminium bromide 34	1.43-6.66	400-409
Stannous chloride	SnCl ₂	189	Pyridine 104	2.5-10.7	824-826
			Ethyl sulphide 104	1.65-4.56	175-176
			$ m Urethane^{136}$	1.36-4.90	183.8-1829

Stannic chloride Stannous bromide Stannic bromide	SnC ₁ SnBr ₂ SnBr,	258·87 278·32 278	Nitrobenzene ¹³⁰ Pyridine ¹⁰⁴ Ethyl sulphide ¹⁰⁴ Bromine ⁴⁹	2.13-6.83 0.66-4.13 1.56-4.28	445'9-290'4 264'1-276'2 283-236
	PbCl ₂ PbCl ₂ Pb (NO ₃) ₂ (CH ₃ CO ₀) ₂ Pb	439 27778 330°35 324°9	Bismuth chloride 95 Pyridine 104 Acetic acid 99	1.88-1035 1.064-1.797 1.27-4.43 2.177-4.878	434-441 275:2-285'8 345'50-363'28 324 328
	AsBr ₃ AsI ₃	315	Nitrobenzene ¹³⁰ Bromine ⁴⁹ Carbon disulphide ¹¹¹	3.39-11-10 0.40-10.45 1.91-5.94 5.65-980	109-194 126'7-175'7 312-320 442-466
Arsenious sulphide Antimony trichloride	As,S ₃ SbCi ₃	246°0 226°6	Fenchone ⁸¹ Sulphur ⁸³ Carbonyl chloride ³⁹ Chloroform ¹¹¹	1.00-3.20 0.67-2.65 0.60-3.04 6.30-11.10	442-457 117'5-131'1 242-255 262-284
Antimony pentachloride Antimony tribromide	SbCl ₅ SbBr ₃	297'3 360	Nitrobenzene ¹³⁰ Carbonyl chloride ³⁹ Chloroform ¹¹¹	2.58-10.65 0.60-3.41 5.11-990	255.6-211'9 314-332 383-420
Antimony pentabromide Antimony triiodide	$_{\mathrm{SbI}_{3}}^{\mathrm{SbBr}_{\mathbf{f}}}$	520 501	Aluminium bromide 84 Bromine 49 Carbon disulphide 111	1.62-12.96 3.40-7.96 2.46-3.54	377-825 532-550 462-502
	Sb.S., BiC.J. BiBr., Bil., CH.COO), Bi	336°0 312°08 448 589 385°5	renchone ⁹⁴ Sulphur ⁸³ Nitrobenzene ¹³⁹ Aluminium bromide ⁸⁴ Fenchone ⁸¹ Acetic acid ⁸⁹	1.81-4.40 0.504-7.095 0.68-3.08 0.89-10.57 0.60-1.29	500-498 198'1-185'4 203'9-326'9 450-1677 593-599
Manganous chloride Ferric chloride	MnCl ₂ FeCl ₃	126	Urethane 136, 140 Acetamide 140 Ether 49 Ethyl alcohol 49	0.72-3.09 0.72-3.09 1.124-3.453 1.53-5.43 2.37-11.57	131'3-141'2 51-54'7 170-185 195-163
	FeCl ₂ CoCl ₂	126°6 130	Fyridine 100, 100 Nitrobenzene 130 Pyridine 104 Urethane 136, 140 Bismuth chloride 95 Quinoline 103	2.63-13.75 1.479-4.739 0.33-0.78 0.48-0.90 0.451-0.688 0.396-2.945	130-150 316'9-190'8 112'3-107'9 256'4-253'8 127'6-124'9 132-146

M.W. (Obs.).	145'17 128'99'123'33 62'4'74'8 202'72-216'56	223-230	225-235 119	109-99 ^{.8} 227-333	69.1-83.1 96.5-879	1.96-80.1	167-152	135.5-189.0	257.4-273	275-369	300-463	26.6-57.9	08.08-03.54	0.111-0./11	194.2-220.0			193.7-152.9	/9.45-/0.12			ő		124.0-126.2
Concentration.	0.50 0.41-1.95 0.481-1.772 0.20-2.99	0.669-4.590	0.790-4.495 69.5 (N)	14.4-73·6 (N) 0.47-3·18	0.335-9'309 4'77-75'0 (N)	10.6-75.7 (N)	14.6-56.6 (N)	0.205-6.232	0.46-1'19 22'7-107'0 (N)	1.87-6.29	0.233-3.575	0.207-0.774	73.51-20.40 (n)	6.66 26.70 (")	0.00-30.50 (")	7.05-43.57 (")	(") 92.77.00 (")	13.46 55.03 (11)	0.617-1.062	3.58-7.11	0.328-3.591	(cryos.) 5.84-36.86 (n)	(ebul.) $40.36 \cdot 17.03(n)$	94.43-37.35 (n)
Solvent.	Piperidine 104 Pyridine 104 Acetamide 140 Pyridine 104	Quinoline ¹⁰³ Quinoline ¹⁰³	Quinoline 103 Sulphur dioxide 66	Sulphur dioxide ⁹⁰ Chloroform ¹¹¹ , ¹¹⁶ , ¹¹⁵	Phenol 101 Sulphur dioxide 66	Sulphur dioxide 66	Sulphur dioxide 66, 69	Phenol 101	Sulphur dioxide 66	Chloroform 111, 115	Bromoform 117	Formamide **	Water 125 Ethyl 2120bol 118	Dinhenylamine 118	Dhonol 118	Acetic acid 118	iso-Amyl alcohol 118	Formic acid 118	Formamide 42	115	Bromoform 117	Water 118 (cryo		Ethyl alcohol 118
M.W. (Calcd.).	218.9	129.6	218·6 67·5	81.6	9.56	109.0	201.0		81.57	109.6									136	137.6				
Formula.	CoBr	NiCl	NiBr ₂ CH ₃ NH ₃ CI	(CH ₃) ₂ NH ₂ Cl	(CH ₃) ₃ NHCI	(CH ₃), NCI (CH ₂),NBr	$(CH_3)_4$ NI		C,H,NH,CI	$(C_2H_5)_2^{\prime}$ NH_2C_1									$(C_2H_5)_2 NH_2NO_3$	$(C_2H_5)_3$ NHCI				
Substance.	Cobalt bromide	Nickel chloride	Methylammonium chloride	Diniculy taining in unit chiginge	Trimethylammonium chloride	Tetramethylammonium bromide	Tetramethylammonium iodide		Ethylammonium chloride	Diethylammonium chloride									Diethylammonium nitrate	Triethylammonium chloride				

			Diphenylamine ¹¹⁸ Phenol ¹¹⁸ Acetic acid ¹¹⁸ <i>iso-</i> Amyl alcohol ¹¹⁸ Formic acid ¹¹⁸	6.03-56.66 (n) 8.92-42.80 (n) 4.37-51.25 (n) 38.56-15.41 (n) 4.55-36.74 (n)	143'0-202'8 148'4-153'5 156'5-229'9 164'7-155'8 93'76-93'80	
Triethylammonium bromide	$(C_2H_5)_3$ NHBr	182	Chloroform 111 Bromoform 117 Water 118	3.74-9.34 0.659-3.166 62.95-18-75 (n)	294-403 328-681 118'4-112'9	
			Etnyl alconol Diphenylamine ¹¹⁸ Phenol ¹¹⁸ Acetic acid ¹¹⁸ iso-Amyl alcohol ¹¹⁸	7.19-34.30 (n) $7.19-35.06 (n)$ $6.64-13.43 (n)$ $3.21-32.82 (n)$ $6.64.3-10.42 (n)$	192'2-171'1 203'9-309'6 217'3-225'0 227'4-350'4 245'5-100'8	
Triethylammonium iodide	(C ₂ H ₅₎₃ NHI	229	Chloroform 111 Bromoform 117 Water 118	4.51-7'90 0'353 52'69-18'69 (n)	498-686 428 144-103.1	
			Ethyl alcohol ¹¹⁸ Diphenylamine Phenol Acetic acid	81.39-23.74 (n) 3.11-33.17 (n) 8.93-29.00 (n) 5.86-13.16 (n)	272.6-250.2 212.6-446.7 2795.301.3 446.6-458.2	
Tetraethylammonium chloride	(C ₂ H _{5),} NCl	165.6	200-Amyl arconol Formic acid Chloroform ¹¹¹ Water ¹¹⁸ Ethyl alcohol ¹¹⁸	44 70-14 40 (n) 7.38 (n) 2.00-6.43 47.97-18.92 (n) 54.54-21.76 (n)		
Tetraethylammonium bromide	$(C_2H_b)_4$ NBr	210'1	Phenol 118 iso-Amyl alcohol 118 Chloroform 111 Water 118 Ethyl alcohol 118 Ethyl alcohol 118	5'73-12'60 (n) 22'13-70'73 (n) 4'26-7'97 56'37-12'69 (n) 39'81-10'66 (n)		
Tetraethylammonium iodide	(C ₂ H ₅)4 NI	257.1	Accent acut Sulphur dioxide 66, 69 Methyl alcohol ¹¹⁹ Ethyl alcohol ¹¹⁹ Accetonitrile ¹¹⁹ Pronionitrile ¹¹⁹	5 30-42 20 (n) 34 49-13 78 (n) 6 41 42 9 (n) 4 383 8 867 1 533 - 3 609 2 2 3 3 - 5 492 1 4 0 2 - 3 001	255 1-560 3 236 3-312 7 236 190 199 0- 208 1 202 7- 230 3 163 1- 175 0 167 0- 182 0	
				•		

		,	Bromoform ¹¹⁷ Ethyl alcohol ¹¹⁸ Phenol ¹¹⁸ Acetic acid ¹¹⁸ <i>iso-</i> Amyl alcohol ¹¹⁸	0'314-1'452 163'6-29'18 (n) 16'80-53'27 (n) 9'59-64'22 (n) 62'63-19'34 (n)	435-466 116'4-110'6 136'0-145'3 155'0-204'2 185'8-173'8 67'61-61'06	
iso-Butylammonium iodide iso-Amylammonium chloride	$(C_5^{}H_9^{})_4^{}NI$ $(C_5^{}H_{11}^{})$ $NH_3^{}CI$	201°1 123°6	Chloroform 111, 115 Chloroform 111, 115 Formamide 42	4.67-9.58 2.77-5.37 0.200-0.634	990-1443 739-703 65:6-60:6	
Di-iso-amylammonium chloride Tetra-iso-amylammonium iodide	e $(C_5H_{11})^2 NH_2C_1 $ $(C_5H_{11})_4 N_1^2$	193.6 427.3	Formamide 42 Chloroform ¹¹¹ Bromoform 117	0.229-0.307 3.15-5.17	528-636 528-636 567-1463	
Aniline hydrochloride Methylaniline hydrochloride	$C_6H_5NH_9$. $HC1$ $C_6H_5NH(CH_3)$. $HC1$	129'6 143'5	Phenol 101 Chloroform 111 Bromoform 117	2.33-5.50	308-351 308-351	
Ethylaniline hydrochloride	$\mathrm{C_6H_5NH(C_2H_5).HC1}$	157.6	Chloroform 111 Water 115	2.40-4.49 2.40-4.49 42.36-19.32	346-353 346-353 100.0-85.1	
			Ethyl alcohol ¹¹⁸ iso-Amyl alcohol ¹¹⁸ Acetic acid ¹¹⁸	96'56-30'31 (n) 50'00-14'07 (n) 4'78-41'21 (n)	192'8-169'2 270'3-320'7 217'4-271'9	
Ethylaniline hydriodide Diethylaniline hydrochloride	$egin{aligned} \mathbf{C_6H_6NH}(\mathbf{C_2H_5}).\mathrm{HI} \\ \mathbf{C_6H_5N}(\mathbf{C_2H_5})_2.\mathrm{HCI} \end{aligned}$	249'I I58'6	Chloroform 111 Chloroform 111 Bromoform 117	3.73-7.51	504-569 256-294	
Diethylaniline hydriodide	$\mathrm{C_6H_5N}(\mathrm{C_2H_5})_2.\mathrm{HI}$	277.0	Chloroform 111 Bromoform 117	5.62-10.25	528-616 528-616	
Benzylammonium chloride Dibenzylammonium chloride Tribenzylammonium chloride	$C_0H_0CH_0NH_3CI \\ (C_0H_0CH_3)_1NH_2CI \\ (C_0H_0CH_2)_3NHCI \\ (C_0H_0CH_2)_3NHCI \\$	143.6 233.5 323.6	Sulphur dioxide 66 Chloroform 111 Chloroform 111	7.87-84.9 (N) 0.48-0.79 2.98-4.68	182-309 487-481 392-397	
Methylpropylphenylbenzylam-	$\mathrm{CH_{3},C_{3}H_{7},C_{6}H_{5},(C_{6}H_{5}\mathrm{CH_{2}})\mathrm{NBr}}$	320	Bromoform 120	0.724	704.8	
monium bromide Methylpropylphenylbenzylam-	CH ₃ ,C ₃ H ₇ ,C ₆ H ₅ ,(C ₆ H ₅ CH ₂)NI	366.8	Bromoform 120	0.286	786	
monium logide Methylallylphenylbenzylam-	CH3.C3H5.C6H5.(C6H5CH2)NBr	318	Bromoform 120	1.692	1.594	
monum bromae Methyliso-butylphenylbenzyl ammonium iodide	$\mathrm{CH_3.C_4H_9.C_6H_5.(C_6H_5\mathrm{CH_2})NI}$	380.8	Bromoform 120	0.295	754	

Concentration. M.W. (Obs.), 0.804 964		% © ∵	1.421-2'435 833-892 1'323-2'194 111-119 8'973 227 0'628-2'180 422-644	1.375-4145 134-166 0.647-2.024 436-685 0.45-1.09 374-384 1.265-2.371 555-502 0.830-3.512 599-817		63 r9.	5.009 3.020-3.351 1.020-3.351 1.020-3.351
Solvent, Bromoform ¹²⁰	Chloroform 111 Chloroform 111 Chloroform 111 Chloroform 111 Vater 118 S1 Rithel alroyol 118	18		Water ¹²¹ Diphenylamine ¹²¹ Chloroform ¹¹⁵ Chloroform ¹¹⁵ Chloroform ¹¹⁵	Alcohol 122 Chloroform 115, 122	Alcohol *** Chloroform 115, 122, 121 A 15ch 2 133	1115
M.W. (Calcd.). 471	165.5 210.0 257.0 285.0	172.6	271	397 292 383 305°5	350	397	328
Formula. CH ₃ .C ₃ H ₅ .C ₆ H ₅ .(C ₆ H ₅ CH ₃)NO ₃ SC ₁₀ H ₁₅ O	$C_0H_7N.HCI$ $C_0H_7N.HBr$ $C_0H_7N.HI$ $C_0H_7NC_2H_5I$	C ₆ H ₆ NH.NHC ₂ H ₆ .HCl	C ₉ H ₂ NCH ₃ I	C,H.,C(C,H,J),N.CH,I C,H.,C.(C,H,J),NHCI C,H.,C.(C,H,J),NHI C,H.,C.(C,H,J),NHI	$C_6H_5C.(C_6H_4)_2\ NCH_3Br$	$C_6H_5C.(C_6H_4)_2$ NCH_3I	$C_6H_5C.(C_6H_4)_2$ N.SCN
Substance. Methylallylphenylbenzylam- C	Quinoline hydrochloride Quinoline hydrobromide Quinoline hydriodide Quinoline ethiodide	Ethylphenylhydrazine hydro- chloride Peridine methiolide	Quinoline methiodide	Phenylacridine methiodide Phenylacridonium chloride Phenylacridonium iodide Methylphenylacridonium chlo- ride	Methylphenylacridonium bro- mide	Methylphenylacridonium iodide	Methylphenylacridonium thio- cyanate

305-421	310 181-260 599-714 949-1359 1765-2951
181,1-692,0	4.83 6.12-55:5 (N) 2.72-5:68 4.02-8:50 5.73-12-48
$Bromoform^{117}$	Chloroform ¹¹¹ Sulphur dioxide ^{66, 69} Chloroform ¹¹¹ Chloroform ¹¹¹ Chloroform ¹¹¹
160.5	204 154'6 199'1 246'1
O HCI	$CH = C$ CH_3 $(CH_3)_3SI$ $(C_2H_5)_3SI$ $(C_2H_5)_3SI$ $(C_2H_5)_3SI$
Dimethylpyrone hydrochloride	Trimethylsulphonium iodide Triethylsulphonium chloride Triethylsulphonium bromide Triethylsulphonium iodide

CH3

cH = c

The following substances or groups of substances have the normal molecular weight in a variety of solvents: sucrose and mannitol 14, 38, 53; halogen derivatives of pseudocuminol and p-xylenol 20; acylcresols 24; halogen derivatives of benzene azophenol 20; pseudocuminolazophenol 20; benzene- and tolueneazo-cresols, xylenols, pseudocuminols and naphthols 20; alkyloxy derivatives of benzene azophenol 20; thiophenol, its halogen and alkyloxy derivatives 11, 1; ethers and esters 1, 8, 12, 14, 19 (except when containing an associating group); camphor 7, 8, 12, 14, 18; benzil 14, 37, 51, 54; quinones 1, 51; secondary and tertiary amines 11, 12, 54, 71, 146; hydrazones 20; azoles and thienyls 12, 51; azobenzene, its amido and oxyderivatives 1, 38; hydrocarbons (see note at head of Appendix) 15, 27, 33, 35, 38, 51, 54, 66, 69, 75, 145, 146; halogen derivatives of hydrocarbons 111, 84; nitro-derivatives of aromatic hydrocarbons (see also note above) 33; nitroso-benzene and toluene and their halogen derivatives 24, 132; iodine 8, 145, 147; organic solutes dissolved in salt hydrates 148.

ADDENDUM.

Published too late for inclusion in the above tables may be mentioned the determinations of the degree of association in nitrobenzene of phenols, alcohols, aromatic and aliphatic acids, aldehydes, ketones, acid anhydrides, amines, amides, anilides, methane and benzyl nitrile by Böeseken and van der Eerden ¹⁵³.

Note on Molecular Weights of Pure Liquids.

The molecular weights of substances in the liquid condition so far determined are of doubtful value. An excellent summary of measurements depending on surface tension measurements is given in the publication by Walden and Swinne (loc. cit., p. 115). In addition may be mentioned an investigation by Walden of the molecular magnitudes of molten organic ammonium salts. (Bull. Acad. Sci. St. Pétersbourg, 1914, p. 405.)

December, 1914.

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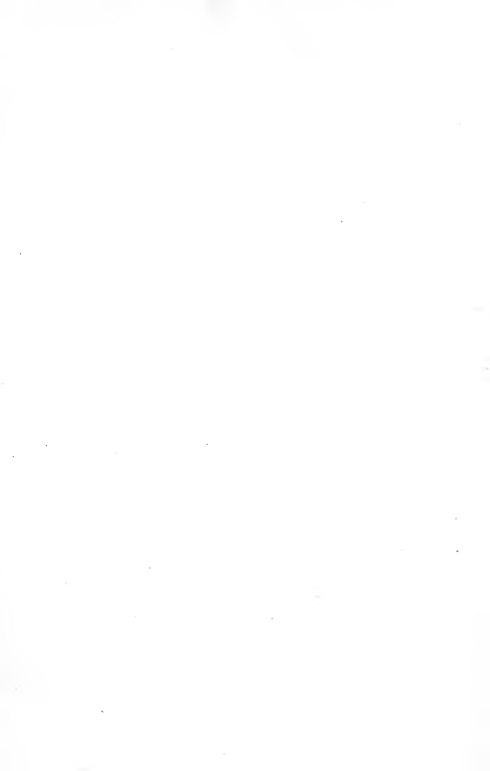
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INDEX

ALLOTROPY, dynamic theory of, 5, 92-5. Aqueous solution, association in, 44, 45. Aromatic acids, behaviour in solution, 33, 35, 36.

Associated substances, Auwers' classes, 33.

- mutual action of, 48, 108.

Association, and chemical combination, 105, 106.

and electro-negative groups, 25, 41, 42, 84, 109.

definition of, 1.

- limits of, 30, 34.

BATSCHINSKI'S method, 65, 84, 85, 86. Bennett and Mitchell's method, 67, 84, 85, 86. Bingham's method, 76.

Boiling point and association, 51, 74.

CHEMICAL action, association theory of, 105.

effect on molecular weight results, 23, 33, 37. Co-efficient of molecular surface energy,

additive character, 62, 63, 68.

Colloids, molecular size of, 30.

Concentration, effect on molecular complexity, 33, 86, 98.

Condensation, molecular change on, 87, 88.

Contraction on solution, effect of association, 77.

Co-volume method, 77, 78.

Critical co-efficient, 54, 55, 100.

density, 66.

pressure, 54, 65.

– temperature, 54, 56, 63, 64, 65, 66.

– volume, 53, 54, 65.

Crystal form and molecular complexity, 90, 91, 94.

DIELECTRIC constant, 31, 41-3, 46-8, 54, 90, 98, 100, 103, 107, 109. Dihydrol, 90, 91.

Dissociation, 1.

Dutoit and Moijou's method, 70-2, 81, 84,

Dynamic allotropy, 5, 92-5.

ELECTRICAL conductivity, 102, 103.

Electrolytes, association of, 3, 25, 27, 28, 32, 42, 46-8, 59, 75, 76, 102, 103.

Electrolytic dissociation hypothesis, 5, 48.

Electro-negative groups, effect of, 25, 41, 42, 84, 109.

FLUIDITY and degree of complexity, 76,

Freezing point depression constant, effect on association, 41.

Guye's methods, 54, 79, 84, 85, 86, 93.

HEAT of association, 17, 32, 84. - dilution, effect of, 38.

Holmes' method, 81. Hydrol, 90, 91.

Hydronal, 91.

Hydrone, 91.

ICE, molecular constitution, 87, 90, 91. Ionizing power and association, 103.

LATENT heat, of evaporation, 21, 49, 51-3, 70, 100.

— — of fusion, 21, 75, 90, 93, 100. Longinescu's method, 74, 84, 85, 86.

MELTING point, a corresponding temperature, 72, 73, 75.

- of solvent, effect on association, 41. Metacritical temperature, 66.

Molecular cohesion, 62, 71, 75.

compounds, 106, 107.

- formulæ, choice of, 14, 96-9.

refraction, 54.

- surface energy, abnormal temperature coefficients, 61, 62.

and substitution, 62.

 weight, determination in solution, 21, 22, III.

Monatomicity, cases of, 10, 11, 17, 23, 29. Morgan's method, 64.

NERNST-THOMSON hypothesis, 42. Normal liquids, true molecular weight of, 49, 86, 87.

OPTICAL rotation and association, 103.

Organic compounds, association of, 25, 26.

Periodic table and association, 12, 23, 25,

Pressure, effect on molecular complexity, 17, 89, 98, 109.

RAMSAY and Shields' method, 57-64, 65, 66, 67, 68, 71, 72, 76, 84, 85, 86. Rectilinear diameter, 53, 54.

Solid solution, effect on molecular weight determination, 23, 24, 40.

molecular weight in, 28, 29, 95. Solubility and association, 82, 101, 102. Solvent, action of solute on, 45, 91, 95. - effect on complexity of solute, 28, 31, 37-48.

Specific cohesion, 57, 64, 65, 69, 70, 72,

73, 85. - heat, 100.

- heats of gases, ratio of, 6, 10.

- inductive capacity, 42. Spiral manometer, 3, 6, 7. Surface energy, 94, 100. - tension, 57, 64, 67, 100.

Partition law, application of, 21, 28, 40, Surface tension methods, objections to, 59, 60, 61, 62, 63, 85, 86.

> TEMPERATURE, effect on molecular complexity, 1, 2, 13, 14, 28, 31, 32, 89, 98, rog. Total surface energy, 67, 68.

Traube's methods, 77, 78, 84, 85, 86. Trihydrol, 90, 91.

Trouton's Rule, 49, 52, 53, 63, 70, 75, 76, 84.

UNSATURATION and association, 109.

VAN DER WAALS' equation, application of, 79, 81, 87, 88. Vapour density, determination of, 7-10. - pressure and association, 50. Velocity of sound method, 10. Viscosity and association, 55, 90, 100. - relationships, 66, 76. Volatile solutes, 40.

WALDEN'S methods, 70, 71, 72, 75, 81, 84, 85. Walden-Kistiakowsky method, 70, 71, 72. Weight drop method, 64.







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